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                THREE NEW FIELDS ADDED TO IFIPAT/IFIUDB/IFICDB
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                (ROSPATENT) added to list of core patent offices covered
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NEWS 19 FEB 16 STN User Update to be held in conjunction with the 229th ACS
                National Meeting on March 13, 2005
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                data from INPADOC
NEWS 21 FEB 28 BABS - Current-awareness alerts (SDIs) available
NEWS 22 FEB 28 MEDLINE/LMEDLINE reloaded
NEWS 23 MAR 02 GBFULL: New full-text patent database on STN
NEWS 24 MAR 03
                REGISTRY/ZREGISTRY - Sequence annotations enhanced
NEWS 25 MAR 03 MEDLINE file segment of TOXCENTER reloaded
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L5 ANSWER 1 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2005:228330 CAPLUS

TITLE:

Apigenin inhibits VEGF and HIF-1 expression via

PI3K/AKT/p70s6K1 and HDM2/p53 pathways

AUTHOR(S):

Fang, Jing; Xia, Chang; Cao, Zongxian; Zheng, Jenny

Z.; Reed, Eddie; Jiang, Bing-Hua

CORPORATE SOURCE:

The Mary Babb Randolph Cancer Center, Department of Microbiology, Immunology and Cell Biology, West

Virginia University, Morgantown, WV, 26506-9300, USA

SOURCE:

FASEB Journal (2005), 19(3), 342-353 CODEN: FAJOEC; ISSN: 0892-6638

PUBLISHER:

Federation of American Societies for Experimental

Biology

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB Apigenin is a nontoxic dietary flavonoid that has been shown to possess anti-tumor properties and therefore poses special interest for the development of a novel chemopreventive and/or chemotherapeutic agent for cancer. Ovarian cancer is one of the most common causes of cancer death among women. Here we demonstrate that apigenin inhibits expression of vascular endothelial growth factor (VEGF) in human ovarian cancer cells. VEGF plays an important role in tumor angiogenesis and growth. We found that apigenin inhibited VEGF expression at the transcriptional level through expression of hypoxia-inducible factor lot (HIF-lot). Apigenin inhibited expression of HIF-lot and VEGF via the PI3K/AKT/p70S6K1 and HDM2/p53 pathways. Apigenin inhibited tube formation in vitro by endothelial cells. These findings reveal a novel role of apigenin in inhibiting HIF-1 and VEGF expression that is important for tumor angiogenesis and growth, identifying new signaling mols. that mediate this regulation.

5 ANSWER 2 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2005:15059 CAPLUS

DOCUMENT NUMBER:

142:169994

TITLE:

Curcumin induces glutathione biosynthesis and inhibits

NF-κB activation and interleukin-8 release in

alveolar epithelial cells: Mechanism of free radical

scavenging activity

AUTHOR (S):

Biswas, Saibal K.; McClure, Danny; Jimenez, Luis A.;

Megson, Ian L.; Rahman, Irfan

CORPORATE SOURCE:

Centre for Cardiovascular Sciences, School of

Biomedical and Clinical Laboratory Sciences, Medical

School, University of Edinburgh, Edinburgh, UK

SOURCE:

Antioxidants & Redox Signaling (2005), 7(1 & 2), 32-41

CODEN: ARSIF2; ISSN: 1523-0864

PUBLISHER: Mary Ann Liebert, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

AB Oxidants and tumor necrosis factor- α (TNF- α) activate

transcription factors such as nuclear factor-κB (NF-κB), which

is involved in the transcription of proinflammatory mediators, including

interleuking 8 (IL-8). Curcumin (diferulo, lmethane) is a naturall, occurring flavonoid present in the spice turmeric, which has a

long traditional use as a chemotherapeutic agent for

many diseases. We hypothesize that curcumin may possess both antioxidant and antiinflammatory properties by increasing the glutathione levels and

inhibiting oxidant- and cytokine-induced NF- κB activation and IL-8

release from cultured alveolar epithelial cells (A549). Treatment of A549 cells with hydrogen peroxide (H2O2; 100 μ M) and TNF- α (10 ng/mL)

significantly increased NF- κ B and activator protein-1 (AP-1)

activation, as well as IL-8 release. Curcumin inhibited both H202- and

 $TNF\text{-}\alpha\text{-mediated}$ activation of NF- κB and AP-1, and IL-8 release.

Furthermore, an increased level of GSH and glutamylcysteine ligase catalytic subunit mRNA expression was observed in curcumin-treated cells as

compared with untreated cells. Curcumin interacted directly with

superoxide anion (02.-) and hydroxyl radical (.OH) as shown by ESR, quenching the interaction of the radicals with the spin trap, Tempone-H.

This suggests that curcumin has multiple properties: as an oxygen radical scavenger, antioxidant through modulation of glutathione levels, and

antiinflammatory agent through inhibition of IL-8 release in lung cells.

REFERENCE COUNT: 31 THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 3 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:578807 CAPLUS

DOCUMENT NUMBER: 142:148584

TITLE: The flavonoid effect against vinblastine,

cyclophosphamide and paracetamol toxicity by

inhibition of lipid-peroxidation and increasing liver

glutathione concentration

AUTHOR(S): Lahouel, M.; Boulkour, S.; Sequeni, N.; Fillastre, J.

Р.

CORPORATE SOURCE: Laboratoire de Pharmacologie et Phytochimie,

Departement de Biologie, Universite de Jijel, Jijel,

18000, Algeria

SOURCE: Pathologie Biologie (2004), 52(6), 314-322

CODEN: PTBIAN; ISSN: 0369-8114

PUBLISHER: Editions Scientifiques et Medicales Elsevier

DOCUMENT TYPE: Journal LANGUAGE: French

AB The paracetamol and cyclophosphamide are metabolized in the liver by the cytochrome P 450. The formed reactive intermediates are responsible of a hepatocyte depletion of the glutathione and a lipoperoxidn. the

vinblastine is also a chemotherapeutic agent

hepatotoxic and hepatotoxic. Otherwise, **flavonoids** are polyphenols substances of plant origin having some biol. and anti-oxidative properties. However no information is available on their

effects on glutathione and glutathione-s-transferases. In our research, we valued the effect of oral administration of **flavonoids** (diosmine and quercetine) under shape of propolis extract to 60 mg/kg daily during 14 days, on hematol. and hepatic toxicity of a single dose of cyclophosphamide 80 mg/kg by i.v. way, vinblastine 2 mg/kg by i.v. way and the hepatic toxicity of the paracetamol managed by oral way to 200 mg/kg corresponding to 2/3 the DL50 at the rat female albinos wistar. We did a blood numeration, an assessment of serum activities of transaminases and alkali phosphatases as well as quantification of the glutathione and the malondialdehyde (MDA) in liver homogenates of rats treated. Analyses are done at regular intervals; 1, 3, 7 and 14 days after the administration of drugs. In the group of rats treated by the cyclophosphamide paracetamol

alone we observed since the 1st day, an increase of lipid peroxide (MDA) of 120% and a downfall of hepatic glutathione including the group receiving the vinblastine (until 210% of reduction). In the same way a severe leucopenia and a thrombopenia (70% of reduction) are observed between the 3rd

and

the 14th day at rats treated by the chemotherapeutic agents alone (cyclophosphamide and timblastine). The combination of flavoroids with drugs have clearly reduced the effect of drugs toxicity. Indeed, the aphasic observed with the vinblastine, as well as the leucopenia and thrombopenia of the cyclophosphamide are corrected entirely. In the same way, we noted a restoration of rates of peroxide and glutathione. Flavonoids seem to act by activation of the turn over of the glutathione and enzymes stimulating particularly glutathione-s-transferases permitting the captation of the reactive metabolites of the studied drugs.

REFERENCE COUNT: 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 4 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:760087 CAPLUS

DOCUMENT NUMBER: 142:86047

TITLE: Role of flavonoids in the prevention of

haematotoxicity due to chemotherapeutic

agents

AUTHOR(S): Lahouel, Mesbah; Fillastre, Jean Paul

CORPORATE SOURCE: Laboratory of Pharmacology and Phytochemistry,

Department of Biology, University of Jijel, Jijel,

18000, Algeria

SOURCE: Haema (2004), 7(3), 313-320

CODEN: HAGAB8; ISSN: 1108-2682

PUBLISHER: Epsilon
DOCUMENT TYPE: Journal
LANGUAGE: English

Flavonoids are polyphenols widely distributed and known to possess biol. and pharmacol. activities, including anti-inflammatory action against free radicals. Haematotoxicity is the main side-effect of chemotherapeutic agents. Therefore, the protection of chemotherapy toxicity by flavonoids is a new field in tumor therapy. Our study shows that oral administration of 100 mg/kg/daily over two weeks of flavonoids (diosmin, hesperidin, quercetin extracted from propolis and daflon) before chemotherapy injection offers some protection against the haematotoxicity of doxorubicin (DOX), cyclophosphamide (CPM) or daunorubicin (DNR). Female wistar rats were injected with a single dose of 10 mg/kg ADR, 40 mg/kg DNR or were given 100 mg/kg CPM in a single dose. A second group received flavonoids 100 mg/kg/daily before chemotherapy for two weeks. Blood samples were taken at different times: 3, 7, 14 and 28 days after the administration of chemotherapeutic agents. A haematol. depletion was observed following treatment with all chemotherapy agents alone, in the first group of rats. The leukopoenia reached the level of 1.500 cells/µl on day 2, and anemia presented three weeks after treatment. A significant protection of chemotherapy haematotoxicity occurred after pre-treatment with flavonoids 100 mg/kg in all groups. We observed no significant difference between rats receiving the combination of flavonoids and chemotherapy and control group. These results suggest that flavonoids seem to offer protection against chemotherapy toxicity.

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 5 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:1008726 CAPLUS

DOCUMENT NUMBER: 142:68564

TITLE: Xanthohumol, a novel anti-HIV-1 agent purified from

hops Humulus lupulus

AUTHOR (S): Wang, Qian; Ding, Zhi-Hui; Liu, Ji-Kai; Zheng,

Yong-Tang

CORPORATE SOURCE: Laboratory of Molecular Immunopharmacology, Kunming

Institute of Zoology, Graduate School, Chinese Academy

of Sciences, Kunming, 650223, Peop. Rep. China

Antiviral Research (2004), 04(3), 100 104

CODEN: ARSPDR; ISSN: 0166-3542

Elsevier B.V. PUBLISHER:

DOCUMENT TYPE: Journal LANGUAGE: English

SCURCE.

Xanthohumol, prenylchacone flavonoid, is a natural product with multi-biofunctions purified from Hops Humulus lupulus. Its anti-HIV-1 activity was tested in the present study. Results showed that xanthohumol inhibited HIV-1 induced cytopathic effects, the production of viral p24 antigen and reverse transcriptase in C8166 lymphocytes at non-cytotoxic concentration The EC50 values were 0.82, 1.28, and 0.50 μ g/mL, resp. The therapeutic index (TI) was about 10.8. Xanthohumol also inhibited HIV-1 replication in PBMC with EC50 value of 20.74 μg/mL. The activity of recombinant HIV-1 reverse transcriptase and the HIV-1 entry were not inhibited by xanthohumol. The results from this study suggested that xanthohumol is effective against HIV-1 and might serve as an interesting lead compound It may represent a novel chemotherapeutic agent for HIV-1 infection. However, the mechanism of its

anti-HIV-1 effect needs to be further clarified.

REFERENCE COUNT: 21 THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

2004:211283 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 141:235838

TITLE: Protective effect of flavonoids against the

toxicity of vinblastine, cyclophosphamide and

paracetamol by inhibition of lipid-peroxidation and

increase of liver glutathion

Lahouel, Mesbah; Boulkour, Soraya; Segueni, Narimane; AUTHOR (S):

Fillastre, Jean Paul

CORPORATE SOURCE: Laboratory of Pharmacology and Phytochemistry,

Department of Biology, University of Jijel, Jijel,

18000, Algeria

SOURCE: Haema (2004), 7(1), 59-67

CODEN: HAGAB8; ISSN: 1108-2682

PUBLISHER: Epsilon DOCUMENT TYPE: Journal LANGUAGE: English

Paracetamol and cyclophosphamide are metabolized in the liver by the AB cytochrome P 450. The produced reactive intermediates are responsible for hepatocyte depletion of glutathione and for lipoperoxidn. Vinblastine is a chemotherapeutic agent, which is also hepatotoxic and hematotoxic. Flavonoids are polyphenols, substances of plant origin, having biol and anti-oxidative properties. There is no available information for the effect of flavonoids on glutathione and glutathione-s-transferases. In our research, we evaluated the effect of oral administration of flavonoids (diosmine and quercetine; 60 mg/kg daily for 14 days) on hematol. and hepatic toxicity of a single dose of cyclophosphamide (80 mg/kg, IV), and vinblastine (2 mg/kg, IV), as well as on the hepatic toxicity of paracetamol given to We measured a full blood count, serum levels of transaminases and alkaline phosphatase as well as levels of glutathione and malondialdehyde (MDA) in liver homogenates of the rats treated. Analyses were performed at regular intervals: 1, 3, 7 and 14 days after the administration of drugs. In the group of rats treated by cyclophosphamide or paracetamol alone, an increase of lipid peroxide (MDA) of 120% and a reduction of hepatic glutathione were observed These changes started after the first day of

treatment. Severe leucopenia and thrombocytopenia (70% of reduction) were also observed between the 3rd and the 14th day in rats treated with the chemotherapeutic agents alone (cyclophosphamide or vinblastine). The combination of flavonoids and chemotherapy produced a clear reduction of drugs toxicity. Bone marrow aplasia, leucopenia and thrombocytopenia were corrected entirely. Furthermore, a restoration of posexide and glutathione was also observed Flavonoids seem to act by the activation of glutathione turnover and enzymes that stimulate

REFERENCE COUNT: 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

particularly glutathione-s-transferases, which permit the capitation of

ANSWER 7 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

the reactive metabolites of the studied drugs.

2004:486847 CAPLUS ACCESSION NUMBER:

141:46900 DOCUMENT NUMBER:

TITLE: Inhibition of cyclooxygenase-2 activity in head and

neck cancer cells by genistein

AUTHOR (S): Ye, Fei; Wu, Josephine; Dunn, Trish; Yi, Jizu; Tong,

Xiaodi; Zhang, David

Department of Pathology, Mount Sinai School of CORPORATE SOURCE:

Medicine, New York University, New York, NY, 10029NY,

Cancer Letters (Amsterdam, Netherlands) (2004), SOURCE:

211(1), 39-46

CODEN: CALEDQ; ISSN: 0304-3835

Elsevier PUBLISHER: DOCUMENT TYPE: Journal English LANGUAGE:

Genistein, rich in soybean, has been reported to have anti-cancer activity on several cancers. However, the mol. mechanism of its anti-cancer activity still remains unclear. We investigated the effect of genistein on a human oral squamous carcinoma line (SCC-25), and demonstrated that qenistein inhibited SCC-25 cell growth via G2/M phase arrest. We observed a significant decrease of proliferating cell nuclear antigen expression in these cells after treatment, but no significant change in the number of apoptotic cells, indicating that the major action of genistein is inhibition of cancer cell proliferation. We also observed a high level of prostaglandin E2 (PGE2) in these cells and PGE2 synthesis in SCC-25 cells was significantly suppressed by genistein. We demonstrated that genistein directly inhibited cyclooxygenase-2 (COX-2) activity, an inducible enzyme that converts arachidonic acid to prostaglandins, similar to the action of celecoxib, a selective COX-2 inhibitor. However, the anticancer activity of genistein was much weaker than that of indomethacin (non-selective COX inhibitor), celecoxib and baicalein (flavonoid isolated from Scutellaria baicalensis). These results suggested that genistein might be useful as a chemopreventive agent rather than a chemotherapeutic

agent.

REFERENCE COUNT: 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

2003:557093 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 139:390885

TITLE: Inhibition of Cancer Cell Proliferation and

Prostaglandin E2 Synthesis by Scutellaria Baicalensis

Zhang, David Y.; Wu, Josephine; Ye, Fei; Xue, Li; AUTHOR (S):

Jiang, Shiquan; Yi, Jizu; Zhang, Wandi; Wei, Huachen;

Sung, Max; Wang, Wayne; Li, Xiaoping

CORPORATE SOURCE: Department of Pathology, Mount Sinai School of

Medicine, New York, NY, 10029, USA

SOURCE: Cancer Research (2003), 63(14), 4037-4043

CODEN: CNREA8; ISSN: 0008-5472

PUBLISHER: American Association for Cancer Research DOCUMENT TYPE: Journal LANGUAGE: English

Scutellaria baicalensis is a widely used Chinese herbal medicine that has been used historically in anti-inflammatory and anticancer therapy. purpose of this study is to verify its anticancer activity on head and neck squamous cell carcinoma (HNSCC) in vitro and in vivo and to investigace its effect on cyclooxygenase 2 (COX 2), which converts arachidonic acid to prostaglardin E2 (PGE2) and is highly expressed in HNSCC. Two human HNSCC cell lines (SCC-25 and KB) and a nontumorigenic cell line (HaCaT) were tested in vitro for growth inhibition, proliferation cell nuclear antigen expression, and COX-2 activity and expression after treatment with Scutellaria baicalensis extract Its effects were compared with those of baicalein (a flavonoid isolated from Scutellaria baicalensis), indomethacin (a nonselective COX inhibitor), and celecoxib (a selective COX-2 inhibitor). Four nude mice with s.c. inoculation of KB cells were tested for its anticancer activity in vivo by oral administration of Scutellaria baicalensis at a dose of 1.5 mg/mouse (75 mg/kg), five times/wk for 7 wk. Scutellaria baicalensis and other agents demonstrated a strong growth inhibition in both tested human HNSCC cell lines. No growth inhibition of HaCaT cells was observed with Scutellaria baicalensis. The 1C50s were 150 µg/mL for Scutellaria baicalensis, 25 µM for celecoxib, and 75 µM for baicalein and indomethacin. Scutellaria baicalensis, as well as celecoxib and indomethacin, but not baicalein, suppressed proliferation cell nuclear antigen expression and PGE2 synthesis in both cell types. Scutellaria baicalensis inhibited COX-2 expression, whereas celecoxib inhibited COX-2 activity directly. A 66% reduction in tumor mass was observed in the nude mice.

Scutellaria baicalensis selectively and effectively inhibits cancer cell growth in vitro and in vivo and can be an effective chemotherapeutic agent for HNSCC. Inhibition of PGE2 synthesis via suppression of COX-2 expression may be responsible for its anticancer activity. Differences in biol. effects of Scutellaria baicalensis compared with baicalein suggest the synergistic effects among

components in Scutellaria baicalensis.

REFERENCE COUNT: 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 9 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2003:198491 CAPLUS

DOCUMENT NUMBER: 139:223762

TITLE: Baicalein and baicalin as inhibitors of HIV-1

integrase

AUTHOR(S): Lee, Min Jun; Kim, Mira; Lee, Yong Sup; Shin, Cha-Gyun

CORPORATE SOURCE: Department of Biotechnology, Chung-Ang University,

Ansung, Kyungki, 456-756, S. Korea SOURCE:
Yakhak Hoechi (2003), 47(1), 46-51 CODEN: YAHOA3; ISSN: 0513-4234
PUBLISHER:
Pharmaceutical Society of Korea

DOCUMENT TYPE: Journal LANGUAGE: Korean

AB Baicalein and baicalin are flavonoid compds. isolated from medicinal herb Scutellaria baicalensis Georgi (Labiatae) and have been known to possess antiviral activities. In the present study, we investigated the in vitro effects of baicalein and baicalin on the three distinctive enzymic activities of the human immunodeficiency virus type-1 (HIV-1) integrase - endonucleolytic, integration, and disintegration activities. Both compds. inhibited the three enzymic activities in a dose-dependent manner. The 50% inhibitory concns. of baicalein and baicalin for endonucleolytic activities of HIV-1 integrase were 4.4±3.3 and 25.9±4.0 μM, resp. In general, baicalein exhibited nearly 6- to

10-fold stronger inhibition than baicalin for the three enzymic activities. These data demonstrate that baicalein or baicalin can be used as a leading compound to develop anti-AIDS chemotherapeutic

agents targeting to the HIV-1 integrase.

L5 ANSWER 10 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:526480 CAPLUS

DOCUMENT NUMBER: 138:117325

TITLE: High-throughput measurement of the Tp53 response to

ancionnect drugs and random compounds using a stably

integrated

AUTHOR(S): Sohn, Taylor A.; Bansal, Ravi; Su, Gloria H.; Murphy,

Kathleen M.; Kern, Scott E.

CORPORATE SOURCE: Department of Surgery, The Johns Hopkins Medical

Institutions, Baltimore, MD, USA

SOURCE: Carcinogenesis (2002), 23(6), 949-957

CODEN: CRNGDP; ISSN: 0143-3334

PUBLISHER: Oxford University Press

DOCUMENT TYPE: Journal LANGUAGE: English

Human Tp53 is normally a short-lived protein. Tp53 protein is stabilized and levels are increased in response to a variety of cellular stresses, including those induced by genotoxic anticancer drugs and environmental exposures. To engineer an efficient assay based on this property, we constructed and integrated a Tp53-specific reporter system into human cancer cells, termed p53R cells. We tested a range of conventional chemotherapeutic agents as well as over 16 000 diverse small compds. Ionizing radiation and two-thirds of conventional chemotherapeutic agents, but only 0.2% of diverse compds. activated Tp53 activity by two-fold or greater, consistent with the presumptive genotoxic activation of Tp53 function. Cytotoxicity was independent of TP53 genetic status when paired, syngeneic wild-type TP53 and TP53-null cells in culture were treated with compds. that activated Tp53. From the unbiased survey of random compds., Tp53 activation was strongly induced by an analog of AMSA, an investigational anti-cancer agent. Tp53 was also strongly induced by an N-oxide of quinoline and by dabequine, an exptl. antimalarial evaluated in humans; dabequine was reported to be neg. in other screens of mutagenicity and clastogenicity but carcinogenic in animal studies. Further exploration of antimalarial compds. identified the common medicinals chloroquine, quinacrine, and amodiaquine as Tp53-inducers. Flavonoids are known to have DNA topoisomerase activity, a Tp53-inducing activity that is confirmed in the assay. A reported clin. association of Tp53 immunopos. colorectal cancers with use of the antihypertensive agents was extended by the demonstration of hydralazine and nifedipine as Tp53-inducers. P53R cells represent an efficient Tp53 functional assay to identify chems. and other agents with

agents, as an adjunct in the pharmaceutical optimization of lead compds., in the exploration of environmental exposures, and in chemical probing of the Tp53 pathway.

interesting biol. properties, including genotoxicity. This assay may have

REFERENCE COUNT: 60 THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 11 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

utility in the identification of novel chemotherapeutic

ACCESSION NUMBER: 2002:461862 CAPLUS

DOCUMENT NUMBER: 138:49527

TITLE: Effects of luteolin on the inhibition of proliferation

and induction of apoptosis in human myeloid leukaemia

cells

AUTHOR(S): Ko, W. G.; Kang, T. H.; Lee, S. J.; Kim, Y. C.; Lee,

В. Н.

CORPORATE SOURCE: College of Pharmacy and Medicinal Resources Research

Center, Wonkwang University, Jeonbuk, 570-749, S.

Korea

SOURCE: Phytotherapy Research (2002), 16(3), 295-298

CODEN: PHYREH; ISSN: 0951-418X

PUBLISHER: John Wiley & Sons Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

Luteolin, a flavonoid isolated from the fruit of Vitex

rotundifolia, has been examined with regard to the inhibition of proliferation and induction of apoptosis in human mveloid leukemia HL-60 dells. The concentration regulared for 500 inhibition of the grouth after 36 h

was 15+1.1 uM. The mode of cell death induced by luteolin was found to be apoptosis, as judged by the morphol. alteration of the cells and by the detection of DNA fragmentation using agarose gel electrophoresis. The degree of apoptosis was quantified by a sandwich enzyme immunoassay and flow cytometric anal. These results suggest that luteolin may be used as potential chemopreventive and chemotherapeutic agents.

REFERENCE COUNT: THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS 16 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 12 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

2002:190371 CAPLUS ACCESSION NUMBER:

TITLE: Preclinical and clinical development of cdk inhibitors

Senderowicz, Adrian M. AUTHOR(S):

Molecular Therapeutics Unit, Oral Pharyngeal Cancer CORPORATE SOURCE:

Branch, NIH, Bethesda, MD, 20892-4340, USA

SOURCE: Abstracts of Papers, 223rd ACS National Meeting,

Orlando, FL, United States, April 7-11, 2002 (2002), MEDI-252. American Chemical Society: Washington, D.

C.

CODEN: 69CKQP

DOCUMENT TYPE: Conference; Meeting Abstract

LANGUAGE: English

Flavopiridol (NSC 649890, HMR 1275) is a flavonoid with potent CDK inhibitory activity. In preclin. models of lymphoid and head and neck cancers, flavopiridol induced apoptosis irresp. of the presence of BCL-2 or p53. The first Phase 1 trial of a cdk inhibitor, flavopiridol, has been completed. The main side effects noted were secretory diarrhea and pro-inflammatory syndrome, while antitumor activity was observed in several tumor types. Concns. between 300 and 500 nM-necessary to inhibit CDK and achieve an antiproliferative effect-were achieved safely. Phase 2 trials with infusional flavopiridol and Phase I infusional trials in combination with standard chemotherapeutic agents are being completed. Another cdk modulator, UCN-01 (7-hydroxystaurosporine; NSC 638850), has also been entered in clin. trials. These first two CDK modulators have shown encouraging results in early clin. trials; the best schedule to be administered and best combination strategies are still under investigation.

ANSWER 13 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

2002:725074 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 138:264985

Antioxidant nutrients and adriamycin toxicity TITLE:

AUTHOR (S): Quiles, Jose L.; Huertas, Jesus R.; Battino, Maurizio;

Mataix, Jose; Ramirez-Tortosa, M. Carmen

CORPORATE SOURCE:

Institute of Nutrition and Food Technology, Department of Physiology, University of Granada, Granada, 18071,

Spain_

Toxicology (2002), 180(1), 79-95 SOURCE:

CODEN: TXCYAC; ISSN: 0300-483X

PUBLISHER: Elsevier Science Ltd. Journal; General Review DOCUMENT TYPE:

English

A review. The anthracycline antibiotic adriamycin (doxorubicin) is one of the most effective chemotherapeutic agents against a wide variety of cancers. However, its use is seriously limited by the

development in the heart of acute and chronic toxic effects. Mechanisms of action and toxicity of adriamycin are briefly revised in this review.

Among followed strategies to attenuate adriamycin toxicity are dosage optimization, synthesis and use of analogs or combined therapy with antioxidants. The most promising results come from the combination of the drug delivery together with an antioxidant to reduce oxidative stress. Many antioxidants have been assayed with very different results. Among these mols., metal ions chelators and low-mol.-mass agents that scavenge reactive exygen species and that are synthesized in vivo have been midely However, the present review will be exclusively focused on the antioxidants that are derived from the diet, in particular the role of vitamin E, vitamin C, vitamin A, coenzyme Q, flavenoids.

antioxidant components of virgin olive oil and selenium.

REFERENCE COUNT:

THERE ARE 143 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE **FORMAT**

ANSWER 14 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

CORPORATE SOURCE:

2001:709408 CAPLUS

DOCUMENT NUMBER:

136:95670

TITLE:

Selective Growth-Inhibitory, Cell-Cycle Deregulatory and Apoptotic Response of Apigenin in Normal versus

Human Prostate Carcinoma Cells

AUTHOR (S):

Gupta, Sanjay; Afaq, Farrukh; Mukhtar, Hasan Department of Dermatology, Case Western Reserve

University and Research Institute of University Hospitals of Cleveland, Cleveland, OH, 44106, USA

SOURCE:

Biochemical and Biophysical Research Communications (2001), 287(4), 914-920

CODEN: BBRCA9; ISSN: 0006-291X

PUBLISHER:

Academic Press

DOCUMENT TYPE:

Journal English

LANGUAGE: Agents that are capable of inducing selective apoptosis of cancer cells are receiving considerable attention in developing novel cancer-preventive approaches. In the present study, employing normal human prostate epithelial cells (NHPE), virally transformed normal human prostate epithelial cells (PZ-HPV-7), and human prostate adenocarcinoma (CA-HPV-10) cells, we evaluated the growth-inhibitory effects of apigenin, a flavonoid abundantly present in fruits and vegetables. Apigenin treatment to NHPE and PZ-HPV-7 resulted in almost similar growth inhibitory responses of low magnitude. In sharp contrast, apigenin treatment resulted in a significant decrease in cell viability of CA-HPV-10 cells. Similar selective growth inhibitory effects were also observed for human epidermoid carcinoma A431 cells compared to normal human epidermal keratinocytes. Apigenin treatment resulted in significant apoptosis of CA-HPV-10 cells as evident from (i) DNA ladder assay, (ii) fluorescence microscopy, and (iii) TUNEL assay, whereas the NHPE and PZ-HPV-7 cells did not undergo apoptosis but showed exclusive necrotic staining only at a high dose of 40 µM. Apigenin (1-10 µM) also resulted in a dose-dependent G2-M phase cell cycle arrest of CA-HPV-10 cells but not of PZ-HPV-7 cells. The growth-inhibitory and apoptotic potential of apigenin was also observed in a variety of prostate carcinoma cells representing different stage and androgen responsiveness may be developed as a promising chemopreventive and/or chemotherapeutic agent against prostate cancer.

2001 Academic Press.

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 15 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

2001:216235 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

135:269

TITLE:

Quercetin inhibits the expression and function of the

androgen receptor in LNCaP prostate cancer cells

AUTHOR (S):

Xing, Nianzeng; Chen, Yi; Mitchell, Susan H.; Young,

Charles Y. F.

CORPORATE SOURCE: Department of Urology and Biochemistry and Molecular

Biology, Mayo Graduate School, Mayo Foundation,

Rochester, MN, 55905, USA

SOURCE: Carcinogenesis (2001), 22(3), 409-414

CODEN: CRNGDP; ISSN: 0143-3334

FUELIGHER: Oxford University Press

DOCUMENT TYPE: Journal LANGUAGE: English

The androgen receptor (AR) is involved in the development and progression of prostate cancer. In order to find new compds. that may present novel mechanisms to attenuate the function of AR, we investigated the effect of a natural flavonoid chemical, quercetin, on androgen action in an androgen-responsive LNCaP prostate cancer cell line. Western blot anal. showed that AR protein expression was inhibited by quercetin in a dose-dependent manner. To demonstrate that the repression effects on AR expression can actually reduce its function, we found that quercetin inhibited the secretion of the prostate-specific, androgen-regulated tumor markers, PSA and hK2. The mRNA levels of androgen-regulated genes such as PSA, NKX3.1 as well as ornithine decarboxylase (ODC) were down-regulated by quercetin. Transient transfections further showed that quercetin inhibited AR-mediated PSA expression at the transcription level. Finally, it was demonstrated that quercetin could repress the expression of the AR gene at the transcription level. Our result suggests that quercetin can attenuate the function of AR by repressing its expression and has the potential to become a chemopreventive and/or chemotherapeutic agent for prostate cancer.

REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 16 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:144762 CAPLUS

DOCUMENT NUMBER: 132:193252

TITLE: Activation and protection of T-cells (CD4+ and CD8+)

using an H2 receptor agonist and other T-cell

activating agents

INVENTOR(S): Hellstrand, Kristoffer; Hermodsson, Svante; Gehlsen,

Kurt R.

PATENT ASSIGNEE(S): Maxim Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 38 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PAT	CENT I	NO.			KINI)	DATE		i	APPL	ICAT:	I NOI	10.		DA	ATE	
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WO	2000	01060	00		A2		2000	0302	1	NO 19	999-t	JS192	211		19	9908	324
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		CZ,	CZ,	DE,	DE,	DK,	DK,	DM,	EE,	EE,	ES,	FI,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,
		SD,	SE,	SG,	SI,	SK,	SK,	SL,	TJ,	TM,	TR,	TT,	UA,	UG,	UZ,	VN,	YU,
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CA	2341	742			AA		2000	0302	(CA 19	999-2	2341	742		19	99908	324
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TW 576745	В	20040221	TW 1999-88114376	;	19990922
AU 9956870	A1	20000314	AU 1999-56870		19990924
AU 765625	B2	20030925			
ZA 2001001787	Α	20010927	ZA 2001-1787		20010302
US 2003039628	A1	20030227	US 2002-265521		20021003
PRIORITY APPLN. INFO.:			US 1998-139281	Α	19980824
			NO 1999-US19211	15	19990824

The present invention relates to a method for facilitating activation of T-cells in a patient, comprising: identifying a patient in need of enhanced T-cell activity, administering an effective amount of a T-cell activating composition to the patient, and administering an effective amount of a

compound that inhibits the production or release of intercellular reactive oxygen metabolites (ROM) to the patient. The present invention further relates to the use of H2-receptor agonists to augment the effectiveness of vaccines. The vaccine composition may also comprises chemotherapeutic agent and/or antiviral agent.

L5 ANSWER 17 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:29

2000:296079 CAPLUS

DOCUMENT NUMBER:

133:4163

TITLE:

Dietary bioflavonoids induce cleavage in the MLL gene

and may contribute to infant leukemia

AUTHOR (S):

Strick, Reiner; Strissel, Pamela L.; Borgers, Susanne;

Smith, Steve L.; Rowley, Janet D.

CORPORATE SOURCE:

Department of Medicine, Section of

Hematology/Oncology, University of Chicago, Chicago,

IL, 60637, USA

SOURCE:

Proceedings of the National Academy of Sciences of the

United States of America (2000), 97(9), 4790-4795

CODEN: PNASA6; ISSN: 0027-8424

PUBLISHER:

National Academy of Sciences

DOCUMENT TYPE: LANGUAGE: Journal English

Chromosomal translocations involving the MLL gene occur in .apprx.80% of infant leukemia. The search for possible agents inducing infant leukemia identified bioflavonoids, natural substances in food and dietary supplements, that cause site-specific DNA cleavage in the MLL breakpoint cluster region (BCR) in vivo. The MLL BCR DNA cleavage was shown in primary progenitor hematopoietic cells from healthy newborns and adults and in cell lines; it colocalized with the MLL BCR cleavage site induced by chemotherapeutic agents, such as etoposide (VP16) and doxorubicin (Dox). Both in vivo and addnl. in vitro expts. demonstrated topoisomerase II (topo II) as the target of bioflavonoids similar to VP16 and Dox. Based on 20 bioflavonoids tested, we identified a common structure essential for the topo II-induced DNA cleavage. Reversibility expts. demonstrated a religation of the bioflavonoid and the VP16-induced MLL cleavage site. The observations support a 2-stage model of cellular processing of topo II inhibitors. The first and reversible stage of topo II-induced DNA cleavage results in DNA repair, but also rarely in chromosome translocations, whereas the second nonreversible stage leads to cell death because of accumulated DNA damage. Thus,

translocations in utero leading to infant and early childhood leukemia.

REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

maternal ingestion of bioflavonoids may induce MLL breaks and potentially

L5 ANSWER 18 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2000:759656 CAPLUS

DOCUMENT NUMBER:

134:13161

TITLE:

Polymethoxyflavonoids from Vitex rotundifolia inhibit proliferation by inducing apoptosis in human myeloid

leukemia cells

AUTHOR(S): Ko, W. G

Ko, W. G.; Kang, T. H.; Lee, S. J.; Kim, N. Y.; Kim,

Y. C.; Sohn, D. H.; Lee, B. H.

College of Pharmacy and Medicinal Resource Research CORPORATE SOURCE:

Center, Wonkwang University, Chonbuk, 570-749, S.

Korea

Food and Chemical Toxicology (2000), 38(10), 861-865 SOURCE:

CODEN: FCTOD7; ISSN: 0278-6915

PUBLISHER: Elsevier Science Led.

POCUMENT TYPE: Journal LANGUAGE: English

Three polymethoxyflavonoids from the fruit of Vitex rotundifolia, namely 2',3',5-trihydroxy-3,6,7-trimethoxyflavone (Vx-1), vitexicarpin (Vx-5) and artemetin (Vx-6), were tested for their antiproliferative activity in human myeloid leukemia HL-60 cells. They showed a dose-dependent decrease in the growth of HL-60 cells. The concns. required for 50% inhibition of the growth (IC50) after 96 h were 4.03 μM , 0.12 μM and 30.98 μM for Vx-1, Vx-5 and Vx-6, resp. Treatment of HL-60 cells with the flavonoids induced morphol. changes that are characteristic of apoptosis. We judged the induction of apoptosis by the detection of DNA fragmentation in agarose gel electrophoresis and the degree of apoptosis was quantified by a double-antibody sandwich ELISA and by flow cytometric The C-3 hydroxyl and C-8 methoxyl groups were found not to be essential for the activity, but the C-3' methoxyl instead of hydroxyl group lowered the antiproliferative and apoptosis inducing activity. These results suggest that the polymethoxyflavonoids isolated from V. rotundifolia may be used as potential chemopreventive and chemotherapeutic agents.

REFERENCE COUNT: THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 19 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:171053 CAPLUS

DOCUMENT NUMBER: 133:53297

TITLE: Application of topoisomerase assays in the evaluation

of natural products as antitumor agents

Constantinou, Andreas; Salti, George AUTHOR(S):

CORPORATE SOURCE: Department of Surgical Oncology, University of

Illinois, Chicago, IL, USA

Journal of Medicinal Food (1999), 2(3-4), 167-171 SOURCE:

CODEN: JMFOFJ; ISSN: 1096-620X

Mary Ann Liebert, Inc. PUBLISHER:

Journal DOCUMENT TYPE: English LANGUAGE:

Initially, DNA topoisomerase (topo) inhibitors found clin. applications as antibiotics and cancer chemotherapeutic agents.

Recently, we demonstrated that plant flavonoids that inhibit

mammalian topo I or topo II might be useful as cancer chemopreventive agents (Constantinou et al., 1995b). Phytochems. can inhibit DNA topoisomerases in different ways; depending on the mode and the type of enzyme, these can be classified as topo I poisons, topo II poisons, topo I antagonists, or topo II antagonists. Correctly classifying topo inhibitors is critical because it provides an important lead as to whether the plant agent can be useful in chemoprevention or in chemotherapy. We outline below a strategy that was designed to identify and classify topo I

and II inhibitors.

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 20 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

1998:258525 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 129:12382

TITLE: Metabolism of the anticancer drug flavopiridol, a new

> inhibitor of cyclin dependent kinases, in rat liver Jager, Walter; Zembsch, Bettina; Wolschann, Peter;

AUTHOR(S): Pittenauer, Ernst; Senderowicz, Adrian M.; Sausville, Edward A.; Sedlacek, Hans H.; Graf, Jurg; Thalhammer,

Therese

CORPORATE SOURCE: Institute of Pharmaceutical Chemistry, University of

Vienna, Vienna, 1090, Austria

SOURCE: Life Sciences (1998), 62(20), 1861-1873

CODEN: LIFSAK; ISSN: 0024-3205

PUBLISHER: Elsevier Science Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

Flavopiridol (I) is a promising novel chemotherapeutic agent currently undergoing clin. phase I trials. The isolated perfused rat liver system was used to examine hepatic metabolism and biliary disposition of I. Besides I two metabolites were detected by high performance liquid chromatog. in bile and perfusate. Twenty-five min after I (30 μ M) addition to the perfusion medium, biliary secretion of metabolite 1 and 2 reached a maximum of 1.04 \pm 0.52 and 11.34 \pm 4.72 nmol/g.liver.min, resp. Biliary excretion of parent I, however, continuously increased for 60 min up to 406 ± 134 pmol/g liver.min. the perfusate, metabolite 1 was below detection limit and release of metabolite 2 was low $(2.8 \pm 0.7 \text{ pmol/g liver.min after 60 min})$. Enzymic hydrolysis with β -glucuronidase, mass spectroscopy and electron absorption spectroscopy revealed that both metabolites are monoglucuronides with the glucuronide in position 5 and 7 of the flavonoid core, resp. The amount of I, metabolite 1 and metabolite 2 excreted into bile during the 60 min of perfusion was 1.94 ± 0.91,

5.15 ± 1.95 and 57.29 ± 23.60 % of I cleared from the perfusate during 60 min, resp. In contrast to the structurally similar flavonoids genistein and daidzein, no inhibition of

UDP-glucuronyltransferase with methylumbelliferone as a substrate was observed indicating that different UDP-glucuronyltransferase isoforms are involved in I metabolism Thus, that glucuronidation is the major mechanism of hepatic I biotransformation. Metabolites are mainly excreted into bile but also released into systemic circulation. The pharmacol. and toxicol. effects of these metabolites remain to be elucidated.

REFERENCE COUNT: 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 21 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:489673 CAPLUS

TITLE: Fractionation of plants to discover substances to

combat cancer. Kinghorn, A. D.

CORPORATE SOURCE: College Pharmacy, University Illinois, Chicago, IL,

60612, USA

SOURCE: Book of Abstracts, 214th ACS National Meeting, Las

Vegas, NV, September 7-11 (1997), AGRO-126. American

Chemical Society: Washington, D. C.

CODEN: 64RNAO

DOCUMENT TYPE: Conference; Meeting Abstract

LANGUAGE: English

AUTHOR(S):

Currently there are four classes of plant-derived drugs used as cancer chemotherapeutic agents in the United States, and a number of plant secondary metabolites found in the human diet are of interest as potential cancer chemotherapeutic agents. In two sep. colaborative multidisciplinary projects, exts. from predominantly tropical plants have been evaluated for their potential anticancer activity and cancer chemopreventive activity using sep. batteries of cell- and mechanism-based in vitro assays. Activity-guided chromatog. fractionation in the presenter's laboratory in these two cancer-based projects has lead to a wide variety of structurally novel bioactive compds., inclusive of alkaloids, diterpenoids, flavonoids, lignans, stilbenoids, triterpenoids, and withanolides, among others. Details of the approaches taken, and of the bioactive compds. obtained, will be discussed.

ANSWER 22 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

1997:285508 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 126:314740

TITLE: Phytochemicals: a glimpse into their structural and

biological variation

AUTHOR (S): Mbwambo, Zakaria H.; Luyengi, Lumonadio; Kinghorn, A.

Douglas

Department of Medicinal Chemistry and Pharmacognosy, COPPORATE SOUPCE:

College of Pharmacy, Program for Collaborative

Research in the Pharmaceutical Sciences, University of

Illinois at Chicago, Chicago, IL, 60612, USA

SOURCE: International Journal of Pharmacognosy (1996), 34(5),

335-343

CODEN: IJPYEW; ISSN: 0925-1618

PUBLISHER: DOCUMENT TYPE:

Journal; General Review

Swets & Zeitlinger

LANGUAGE: English

A review with 48 refs. Higher plants have afforded a plethora of structurally varied biol. active secondary metabolite organic mols. and accordingly have been subjected to wide exptl. scrutiny in countries all over the world. Here, the promise of just one phytochem. group, the

flavonoids, is focused upon. Compds. of this type which have potential use as cancer chemopreventives, cancer chemotherapeutic

agents, and sucrose substitutes are described.

REFERENCE COUNT: 48 THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 23 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

1995:872809 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

123:329296

TITLE:

Quercetin not only inhibits P-glycoprotein efflux activity but also inhibits CYP3A isoenzymes. Reply to

letter to the editors.

AUTHOR (S): Scambia, G.; Ranelletti, F. O.; Benedetti-Panici, P.;

Vincenzo, R. De; Bonanno, G.; Ferrandina, G.;

Piantelli, M.; Mancuso, S.

Facolta di Medicina e Chirurgia "Agostino Gemelli", CORPORATE SOURCE:

Universita Cattolica del Sacro Cuore, Rome, I-00168,

Italy

Cancer Chemotherapy and Pharmacology (1995), 36(5), SOURCE:

449-50

CODEN: CCPHDZ; ISSN: 0344-5704

PUBLISHER: Springer DOCUMENT TYPE: Journal

LANGUAGE: English

A reply to the letter of M.A. Sarkar (ibid., 448-9, 1995) regarding the possible interaction of quercetin with cytotoxic agents via inhibition of cytochrome P 450 CYP3A. It is granted that therapeutic use of quercetin, particularly in combination with chemotherapeutic agents , should be considered with caution. However, it would be important to evaluate the possibility of reducing the therapeutic doses of cytotoxic

agents in combination with quercetin, given that this flavonoid may increase their bioavailability. The detoxifying activity of quercetin may even prove able to reduce the cytotoxic side effects of chemotherapeutic agents.

ANSWER 24 OF 27 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:155161 CAPLUS

DOCUMENT NUMBER: 124:249399

Natural products with immunomodulating and TITLE:

antineoplastic activity

AUTHOR (S): Franz, G.

Inst. Pharmacy, Univ. Regensburg, Regensburg, 93040, CORPORATE SOURCE:

Germany

SOURCE:

Pharmaceutical and Pharmacological Letters (1995),

5(4), 154-8

CODEN: PPLEE3; ISSN: 0939-9488 Medpharm Scientific Publishers

DOCUMENT TYPE:

Journal: General Review

PUBLISHER: LANGUAGE:

English

A review with 30 refs. Immunoscimulants are compds, leading predominantly to a non specific stimulation of the immunol, defense system of treatment is an attractive alternative to conventional chemotherapy and prophylaxis of infections especially when the hosts defense mechanisms have to be activated under conditions of impaired immune responsiveness. Today different screening methods for the detection of immunostimulating compds. are available which allow a determination of the functional state and the efficacy

of the mononuclear phagocyte system, in vivo and in vitro. These include those that use granulocytes, macrophages, T-lymphocyte populations, lymphocyte mitogenic activity, natural killer cells and complement components as target cells or systems. The list of immunol. active agents currently being investigated is quite extensive and constantly enlarging. Many natural compds. have been tested for this purpose and some have been found to be quite active. Emphasis will be presented upon immunol. active carbohydrates, and flavonoids. Mainly for a series of well defined polysaccharides, immune potentiating activities are well documented and closely related to distinct structural features. These biopolymers can be utilized for adjuvant treatment of cancer patients in cotherapy with otherwise immunosuppressive chemotherapeutic agents. However, even if the effects are obvious and reproducible in different in vitro and in vivo systems, only very little is known concerning the absorption, distribution and pharmacokinetics of these substances.

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NEWS 8 DEC 15 MEDLINE update schedule for December 2004
NEWS 9 DEC 17 ELCOM reloaded; updating to resume; current-awareness
                alerts (SDIs) affected
NEWS 10 DEC 17
                COMPUAB reloaded; updating to resume; current-awareness
                 alerts (SDIs) affected
NEWS 11 DEC 17
                SOLIDSTATE reloaded; updating to resume; current-awareness
                 alerts (SDIs) affected
NEWS 12 DEC 17
                CERAB reloaded; updating to resume; current-awareness
                 alerts (SDIs) affected
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                THREE NEW FIELDS ADDED TO IFIPAT/IFIUDB/IFICDB
NEWS 14 DEC 30 EPFULL: New patent full text database to be available on STN
NEWS 15 DEC 30
                CAPLUS - PATENT COVERAGE EXPANDED
NEWS 16 JAN 03 No connect-hour charges in EPFULL during January and
                 February 2005
                CA/CAPLUS - Russian Agency for Patents and Trademarks
NEWS 17 FEB 25
                 (ROSPATENT) added to list of core patent offices covered
                STN Patent Forums to be held in March 2005
NEWS 18 FEB 10
NEWS 19 FEB 16 STN User Update to be held in conjunction with the 229th ACS
                National Meeting on March 13, 2005
NEWS 20 FEB 28
                PATDPAFULL - New display fields provide for legal status
                data from INPADOC
NEWS 21 FEB 28 BABS - Current-awareness alerts (SDIs) available
NEWS 22 FEB 28 MEDLINE/LMEDLINE reloaded
NEWS 23 MAR 02 GBFULL: New full-text patent database on STN
NEWS 24 MAR 03
                REGISTRY/ZREGISTRY - Sequence annotations enhanced
NEWS 25 MAR 03 MEDLINE file segment of TOXCENTER reloaded
NEWS EXPRESS JANUARY 10 CURRENT WINDOWS VERSION IS V7.01a, CURRENT
             MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 10 JANUARY 2005
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             STN Operating Hours Plus Help Desk Availability
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1 GEMCITABINES

L1 2344 GEMCITABINE

(GEMCITABINE OR GEMCITABINES)

=> s flavonoid?

L2 30430 FLAVONOID?

=> s L1 and L2

L3 4 L1 AND L2

=> d 1-4 L3

L3 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN

AN 2004:41226 CAPLUS

DN 140:105321

TI Methods and compositions relating to isoleucine boroproline compounds

IN Adams, Sharlene; Miller, Glenn T.; Jesson, Michael I.; Jones, Barry

PA Point Therapeutics, Inc., USA

SO PCT Int. Appl., 152 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 2

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2004004658 A2 20040115 WO 2003-US21405 20030709

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     2002:526480 CAPLUS
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     138:117325
    High-throughput measurement of the Tp53 response to anticancer drugs and
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     random compounds using a stably integrated
     Sohn, Taylor A.; Bansal, Ravi; Su, Gloria H.; Murphy, Kathleen M.; Kern,
AU
     Scott E.
     Department of Surgery, The Johns Hopkins Medical Institutions, Baltimore,
CS
     MD, USA
so
     Carcinogenesis (2002), 23(6), 949-957
     CODEN: CRNGDP; ISSN: 0143-3334
PB
     Oxford University Press
DT
     Journal
     English
LA
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     2001:132660 CAPLUS
     135:131573
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     Development of cyclin-dependent kinase modulators as novel therapeutic
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     approaches for hematological malignancies
AU
     Senderowicz, A. M.
CS
     Molecular Therapeutics Unit, Oral and Pharyngeal Cancer Branch, National
     Institute of Dental and Craniofacial Research, National Institutes of
     Health, Bethesda, MD, 20892-4340, USA
SO
     Leukemia (2001), 15(1), 1-9
     CODEN: LEUKED; ISSN: 0887-6924
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     1996:510910 CAPLUS
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     125:185131
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     Drug resistance against gemcitabine and topotecan mediated by
     constitutive hsp70 overexpression in vitro: Implication of quercetin as
     sensitizer in chemotherapy
     Sliutz, G.; Karlseder, J.; Tempfer, C.; Orel, L.; Holzer, G.; Simon, MM
AU
     Medical School, University Vienna, Vienna, 1090, Austria
CS
     British Journal of Cancer (1996), 74(2), 172-177
SO
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AND TECHNOLOGY CORPORATION, AND FACHINFORMATIONSZENTRUM KARLSRUHE

FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Mar 18, 2005 (20050318/UP).

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(FILE 'HOME' ENTERED AT 15:49:39 ON 21 MAR 2005)

FILE 'CAPLUS' ENTERED AT 15:49:46 ON 21 MAR 2005

L1 2344 S GEMCITABINE

L2 30430 S FLAVONOID?

L3 4 S L1 AND L2

FILE 'STNGUIDE' ENTERED AT 15:51:02 ON 21 MAR 2005

=> s circiliol

0 CIRCILIOL

L4 0 CIRCILIOL

=> s ?flavone?

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If you are searching in a field that uses implied proximity, and you used a truncation symbol after a punctuation mark, the system may interpret the truncation symbol as being at the beginning of a term. Implied proximity is used in search fields indexed as single words, for example, the Basic Index.

=> s ?flavone

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L6 0 FLAVONE

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=> s flavone?

L7 0 FLAVONE?

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=> fiel caplus

FIEL IS NOT A RECOGNIZED COMMAND

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FILE COVERS 1907 - 21 Mar 2005 VOL 142 ISS 13 FILE LAST UPDATED: 20 Mar 2005 (20050320/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> s flavone

10179 FLAVONE

9033 FLAVONES

L8 1.5562 FLAVONE

(FLAVONE OR FLAVONES)

=> s ?flavone?

L9 26189 ?FLAVONE?

=> s gemcitabine

2344 GEMCITABINE

1 GEMCITABINES

L10 2344 GEMCITABINE

(GEMCITABINE OR GEMCITABINES)

=> s L9 and L10

L11 7 L9 AND L10

=> d 1-7 L11

L11 ANSWER 1 OF 7 CAPLUS COPYRIGHT 2005 ACS on STN

AN 2004:759839 CAPLUS

DN 141:254551

TI Methods and compositions to determine the chemosensitizing dose of suramin used in combination therapy

IN Au, Jessie L. S.; Wientjes, M. Guillaume

PA USA

SO U.S. Pat. Appl. Publ., 19 pp., Cont.-in-part of Appl. No. PCT/US02/30210. CODEN: USXXCO

DT Patent

LA English

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     2004:41226 CAPLUS
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     140:105321
    Methods and compositions relating to isoleucine boroproline compounds
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     Adams, Sharlene: Miller, Glenn T.; Jesson, Michael I.; Jones, Barry
     Point Therapeutics, Inc., USA
PA
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     PCT Int. Appl., 152 pp.
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     2003:913055 CAPLUS
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     139:399770
     Medical goods comprising heparin or chitosan-based hemocompatible coating
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     Horres, Roland; Linssen, Marita Katharina; Hoffmann, Michael; Faust,
IN
     Volker; Hoffmann, Erika; Di Biase, Donato
PA
     Hemoteq G.m.b.H., Germany
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     PCT Int. Appl., 93 pp.
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     2003:261608 CAPLUS
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     138:265631
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     Methods and compositions to determine the chemosensitizing dose of suramin
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     Au, Jessie L.-S.; Wientjes, M. Guillaume
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     2002:695764 CAPLUS
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     Combination therapy for reduction of toxicity of chemotherapeutic agents
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     Prendergast, Patrick T.
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     PCT Int. Appl., 66 pp.
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     2002:555299 CAPLUS
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     Redox therapy for tumors
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     Hoffman, Arnold
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     PCT Int. Appl., 36 pp.
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     2000:742053 CAPLUS
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     Synthesis, activity and formulations of pharmaceutical compounds for
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     Del Soldato, Piero
IN
PA
     Nicox S.A., Fr.
     PCT Int. Appl., 159 pp.
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specification.
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L13 32 L9 AND L12

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1666 CHEMOTHED V DEILLICS

18613 CHEMOTHERAPEUTIC

(CHEMOTHERAPEUTIC OR CHEMOTHERAPEUTICS)

720621 AGENT

1029855 AGENTS

1459468 AGENT

(AGENT OR AGENTS)

8561 CHEMOTHERAPEUTIC AGENT

(CHEMOTHERAPEUTIC (W) AGENT)

30542 ANTICANCER?

L14 38310 CHEMOTHERAPEUTIC AGENT OR ANTICANCER?

=> s L9 and L14

L15 226 L9 AND L14

=> s L1 and L15

L16 3 L1 AND L15

=> d 1-3 ibib abs

L16 ANSWER 1 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:759839 CAPLUS

DOCUMENT NUMBER:

141:254551

TITLE:

Methods and compositions to determine the

chemosensitizing dose of suramin used in combination

therapy

INVENTOR(S):

Au, Jessie L. S.; Wientjes, M. Guillaume

PATENT ASSIGNEE(S):

USA

SOURCE:

U.S. Pat. Appl. Publ., 19 pp., Cont.-in-part of Appl.

No. PCT/US02/30210.

CODEN: USXXCO
PE: Patent

DOCUMENT TYPE: LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

· PA	TENT	NO.			KIN	D	DATE		;	APPL	ICAT:	ION I	. OV		D	ATE	
US	2004	1809	55		A1	-	2004	0916	1	US 2	004-	8076:	20		2	0040	324
WC	2003	0265	74		A2		2003	0403	1	WO 2	002-1	US30:	210		2	0020	924
WC	2003	0265	74		A3		2004	0415									
	W:	ΑE,	AG,	AL,	AU,	BA,	BB,	BG,	BR,	BZ,	CA,	CN,	CO,	CR,	CU,	CZ,	DM,
		DZ,	EC,	EE,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KΡ,
		KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LV,	MA,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,
		NZ,	OM,	PH,	PL,	RO,	SD,	SG,	SI,	SK,	SL,	TN,	TT,	TZ,	UA,	UG,	US,
		UZ,	VN,	YU,	ZA,	ZM,	ZW										
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		CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG			
PRIORIT	Y APF	LN.	INFO	. :	-				1	US 2	001-	3247	04P]	P 2	0010	924
									1	WO 2	002-1	US30:	210	1	A2 2	0020	924

AB A method for determining a therapeutically effective amount of suramin for administering to a patient, who is to receive a cytotoxic agent, which comprises the steps of determining the circulating suramin concentration in the patient; administering suramin, if required, to establish a low circulating concentration of suramin in the patient of below about 200 µM; and administering the chemotherapeutic agent to the

patient when the low circulating concentration of suramin is present in the patient. Conveniently a nomogram can be constructed for use in clin. settings with the suramin.

L16 ANSWER 2 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2003:261608 CAPLUS

DUCUMENT NUMBER:

130:26.031

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Methods and compositions to determine the

chemosensitizing dose of suramin used in combination

therapy

INVENTOR(S):

Au, Jessie L.-S.; Wientjes, M. Guillaume

PATENT ASSIGNEE(S):

USA

SOURCE:

PCT Int. Appl., 47 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	rent	NO.			KIN)	DATE		i	APPL	ICAT:	ION I	NO.		D	ATE	
WO	2003	0265	74		A2	-	2003	0403	1	WO 2	002-1	US30:	210		2	0020	924
WO	2003	0265	74		A 3		2004	0415									
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		DZ,	EC,	EE,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	ΚP,
		KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LV,	MA,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,
		NZ,	OM,	PH,	PL,	RO,	SD,	SG,	SI,	SK,	SL,	TN,	TT,	TZ,	UA,	ŪĠ,	US,
	UZ, VN, YU RW: GH, GM, KE				ZA,	ZM,	ZW										
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
		KG,	ΚZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
		FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	SK,	TR,	BF,	ВJ,	CF,
·		CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG			
EP	1429	713			A2		2004	0623	1	EP 2	002-	7663	46		2	0020	924
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	·AL,	TR,	BG,	CZ,	EE,	SK		
US	US 2004180955						2004	0916	1	US 2	004-	8076	20		2	0040	324
PRIORIT	PRIORITY APPLN. INFO.:								1	US 2	001-	3247	04P		P 2	0010	924
	IORITY APPLN. INFO.:								1	WO 2	002-1	US30:	210	1	W 2	0020	924

AB A method for determining a therapeutically effective amount of suramin for administering to a patient who is to receive a cytotoxic agent comprises determining the circulating suramin concentration in the patient; administering suramin, if required, to establish a low circulating concentration of suramin

in the

the patient of below about 200 μM ; and administering the chemotherapeutic agent to the patient when the low circulating concentration of suramin is present in the patient. Conveniently a nomogram can be constructed for use in clin. settings with the suramin.

L16 ANSWER 3 OF 3 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:695764 CAPLUS

DOCUMENT NUMBER:

137:210932

TITLE:

Combination therapy for reduction of toxicity of

chemotherapeutic agents
Prendergast, Patrick T.

INVENTOR(S):
PATENT ASSIGNEE(S):

Ire.

SOURCE:

PCT Int. Appl., 66 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE: Er FAMILY ACC. NUM. COUNT: 1

English

DATEME INCOMMETON

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002069949	A2	20020912	WO 2002-IB632	20020305

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A3
     WO 2002069949
                                20030605
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, UE, UZ, VW, YU, ZA, ZW, ZW
         RW: GH, CM, KE, I-S. MW, MZ, SD, SI, SZ, TZ, UG, ZM, ZW, AM, AZ, PY,
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             GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA,
             GN, GQ, GW, ML, MR, NE, SN, TD, TG
     US 2002169140
                        A1 20021114
                                           US 2002-91855
                                                                   20020306
                                            IE 2001-209
                                                               A 20010306
PRIORITY APPLN. INFO.:
     Provided in the present invention are compds. suitable for treating
     neoplasms and tumors, viral, bacterial and parasite infections and
     combination therapy with these agents to lower the adverse side effects.
=> d hist
     (FILE 'HOME' ENTERED AT 15:49:39 ON 21 MAR 2005)
     FILE 'CAPLUS' ENTERED AT 15:49:46 ON 21 MAR 2005
           2344 S GEMCITABINE
L1
          30430 S FLAVONOID?
L2
              4 S L1 AND L2
1.3
     FILE 'STNGUIDE' ENTERED AT 15:51:02 ON 21 MAR 2005
L4
              0 S CIRCILIOL
              0 S ?FLAVONE?
L5
L6
              0 S ?FLAVONE
L7
              0 S FLAVONE?
     FILE 'CAPLUS' ENTERED AT 15:58:45 ON 21 MAR 2005
L8
          15562 S FLAVONE
L9
          26189 S ?FLAVONE?
L10
           2344 S GEMCITABINE
L11
             7 S L9 AND L10
L12
           8929 S CHEMOTHERAPEUTIC AGENT OR ANTICANCER COMPOUND
L13
             32 S L9 AND L12
L14
          38310 S CHEMOTHERAPEUTIC AGENT OR ANTICANCER?
L15
            226 S L9 AND L14
L16
             3 S L1 AND L15
=> s circiliol or cirsiciol
             1 CIRCILIOL
             0 CIRSICIOL
L17
             1 CIRCILIOL OR CIRSICIOL
=> d 117
    ANSWER 1 OF 1 CAPLUS COPYRIGHT 2005 ACS on STN
     1994:431157 CAPLUS
DN
     121:31157
ΤI
     Steroidal compounds from Teucrium chamaedrys subsp. chamaedrys
ΑIJ
     Ulubelen, A.; Topcu, G.; Kaya, U.
     Fac. Pharm., Univ. Istanbul, Istanbul, 34452, Turk.
CS
     Phytochemistry (1994), 36(1), 171-3
     CODEN: PYTCAS; ISSN: 0031-9422
DT
     Journal
LΑ
     English
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=> s circiliol or cirsiliol 1 CIRCILIOL

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L13
             32 S L9 AND L12
        38310 S CHEMOTHERAPEUTIC AGENT OR ANTICANCER?
L14
L15
           226 S L9 AND L14
           3 S L1 AND L15
1 S CIRCILIOL OR CIRSICIOL
L17
           125 S CIRCILIOL OR CIRSILIOL
L18
L13
              1 S L13 AMD L15
              1 S 1418 AND 1412
7.20
               1 S L18 AND L10
L21
=> d 1-32 IBIB abs L13
L13 ANSWER 1 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 2005:53052 CAPLUS
DOCUMENT NUMBER:
                         142:190260
TITLE:
                         Genistein, a soy isoflavone, enhances
                          necrotic-like cell death in a breast cancer cell
                          treated with a chemotherapeutic
                          agent
                          Satoh, Haruna; Nishikawa, Kazuhiro; Suzuki, Kazuyuki;
AUTHOR(S):
                         Asano, Ryuji; Virgona, Nantiga; Ichikawa, Tomio;
                          Hagiwara, Kiyokazu; Yano, Tomohiro
CORPORATE SOURCE:
                        Department of Food Science Research for Health,
                          National Institute of Health and Nutrition, Tokyo,
                          162-8636, Japan
                          Research Communications in Molecular Pathology and
SOURCE:
                          Pharmacology (2003), 113-114, 149-158
                          CODEN: RCMPE6; ISSN: 1078-0297
                          PJD Publications Ltd.
PUBLISHER:
                          Journal
DOCUMENT TYPE:
LANGUAGE:
                          English
     Genistein is a major component of soybean isoflavone and has
     preventive effect against breast cancer. In breast cancer, the
     over-expression of HER-2 contributes to malignant transformation of the
     cancer cells. The present study was undertaken to estimate if genistein could
     act as a useful anti-cancer agent against a breast cancer cell
     over-expressing HER-2 in combination with a conventional chemotherapy
     agent, adriamycin (ADR). Genistein enhanced cytotoxic effect of ADR at
     low doses < IC50 against the human breast cancer cell. The enhancing
     effect was mainly dependent on the elevation of necrotic-like cell death
     but not apoptotic cell death. In conjugation with this event, remarkable
     inactivation of HER-2 and Akt in the breast cancer cell was caused by the
     combination of genistein and ADR. These results suggest that genistein
     enhances necrotic-like cell death of the breast cancer cells through the
     inactivation of HER-2 receptor and Akt in combination with ADR.
REFERENCE COUNT:
                                 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS
                          26
                                 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L13 ANSWER 2 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 2004:958643 CAPLUS
ACCESSION NUMBER:

DOCUMENT NUMBER:

142:173464

TITLE:

Anticancer compounds isolated from leaves of Crinum latifolium, derivatives thereof and anticancer composition containing the same

INVENTOR(S):

PATENT ASSIGNEE(S):

S. Korea

Repub. Korean Kongkae Taeho Kongbo, No pp. given
                          CODEN: KRXXA7
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          Korean
FAMILY ACC. NUM. COUNT: 1
```

PATENT NO. KIND DATE APPLICATION NO. DATE

PATENT INFORMATION:

KR 2003061981 Δ 20030723 KR 2002-2145 20020114 PRIORITY APPLN. INFO.: KR 2002-2145 20020114

Anticancer compds. isolated from leaves of Crinum

latifolium, derivs. thereof and an anticancer composition containing the same are

provided, thereby preventing and treating cancers effectively. Anticancer compds, isolated from leaves of Crimum latifolium include 6,7-dimethoxy-4-(3-methyl-2-butenoyloxymethyl)coumarin of the formula I, and 5,6,3-trihydroxy-7,8,4-trimethoxyflavon of the formula II. Another anticancer compound represented by the formula III is provided, wherein R is C1-C3 alkyl. A process for preparing the compound of the formula III comprises the steps of: (i) condensing 3,4-dioxyphenol and Et 4-chloroacetate by a Pechman method in sulfuric acid to prepare 4-chloromethyl-6,7-dimethoxycoumarin; and (ii) reacting 4-chloromethyl-6,7-dimethoxycoumarin with organic acid in the presence of triethylamine.

LIS ANSWER 3 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:759839 CAPLUS

DOCUMENT NUMBER:

141:254551

TITLE:

Methods and compositions to determine the

chemosensitizing dose of suramin used in combination

INVENTOR(S):

Au, Jessie L. S.; Wientjes, M. Guillaume

PATENT ASSIGNEE(S): USA

SOURCE:

U.S. Pat. Appl. Publ., 19 pp., Cont.-in-part of Appl.

No. PCT/US02/30210.

CODEN: USXXCO

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT	NO.			KIN	D	DATE		7	APPL:	ICAT:	ION I	. 01		D	ATE	
		-`				_									-		
US	2004	1.809	55		A1		2004	0916	1	US 2	004-	8076	20		2	0040	324
WO	2003	0265	74		A2		2003	0403	į	WO 2	002-1	US30:	210		2	0020	924
WO	2003	0265	74		A3		2004	0415			,						
	W:	ΑE,	AG,	AL,	AU,	BA,	BB,	BG,	BR,	ΒZ,	CA,	CN,	CO,	CR,	CU,	CZ,	DM,
		DZ,	EC,	EE,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	ΚP,
		KR,	KΖ,	LC,	LK,	LR,	LS,	LT,	LV,	MA,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,
		NZ,	OM,	PH,	PL,	RO,	SD,	SG,	SI,	SK,	SL,	TN,	TT,	TZ,	UA,	UG,	US,
		UZ,	VN,	YU,	ZA,	ZM,	ZW										
	RW:	GH,	GM,	KΕ,	LS,	MW,	MΣ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	AZ,	BY,
		KG,	KΖ,	MD,	RU,	ΤJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	.DE,	DK,	EE,	ES,
		FI,	FR,	GB,	GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	SK,	TR,	BF,	ВJ,	CF,
		CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG			
PRIORITY	APP	LN.	INFO	.:					1	US 2	001-	3247	04P	:	P 2	0010	924
									1	WO 2	002-1	US30:	210	1	A2 2	0020	924

A method for determining a therapeutically effective amount of suramin for administering to a patient, who is to receive a cytotoxic agent, which comprises the steps of determining the circulating suramin concentration in the patient; administering suramin, if required, to establish a low circulating concentration of suramin in the patient of below about 200 µM; and administering the chemotherapeutic agent to the patient when the low circulating concentration of suramin is present in the patient. Conveniently a nomogram can be constructed for use in clin. settings with the suramin.

L13 ANSWER 4 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:675947 CAPLUS

DOCUMENT NUMBER:

141:325120

TITLE:

Clinical characteristics and pharmacokinetics of

purified soy isoflavones: Multiple-dose

administration to men with prostate neoplasia

AUTHOR(S): Fischer, Leslie; Mahoney, Chrysa; Jeffcoat, A. Robert;

Koch, Matthew A.; Thomas, Brian F.; Valentine, John L.; Stinchcombe, Thomas; Boan, Jarol; Crowell, James

A.; Zeisel, Steven H.

CORPORATE SOURCE: Department of Nutrition, School of Public Health,

School of Medicine, University of North Carolina,

Chapel Hill, NC, 27590-7401, UJA

SOURCE: Nutrition and Cancer (2004), 48(2), 160-170

CODEN: NUCADQ; ISSN: 0163-5581

PUBLISHER: Lawrence Erlbaum Associates, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

A phase I clin. trial was conducted to determine the safety, pharmacokinetic parameters, and efficacy of orally administered isoflavones (genistein and daidzein, potential cancer chemotherapeutic agents) over a 3-mo period in men with prostate neoplasia. Twenty men, ages 40 and above, with stage B, C, or D adenocarcinoma of the prostate were treated with a multiple-dose regimen of a soy isoflavone formulation (delivering approx. 300 or 600 mg/day genistein and half this much daidzein) for 84 days. The delivered dose of isoflavones was more than 10-fold higher than that typically taken by prostate cancer patients. In men with prostate cancer, relatively minor side effects of chronic isoflavone treatment were observed including some estrogenic effects (breast changes, increased frequency of hot flashes). Serum dehydroepiandrosterone was decreased by 31.7% (P = 0.0004) at the end of treatment. Except for those subjects whose prostate-specific antigen (PSA) values were below 0.4 ng/mL, subjects had a history of increasing PSA levels prior to the trial. This increase continued during the trial both while on soy isoflavones and after treatment was discontinued. On average the rate of rise accelerated after soy isoflavones were discontinued, but that difference did not attain statistical significance. Genistein and daidzein were rapidly cleared from plasma and excreted in urine. Pharmacokinetic data for chronic dose administration were similar to single-dose administration for the isoflavones investigated except that we observed slightly

longer circulation time for daidzein.

REFERENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 5 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2004:384103 CAPLUS

DOCUMENT NUMBER: 141:17096

TITLE: Genistein inversely affects tubulin-binding

agent-induced apoptosis in human breast cancer cells

AUTHOR(S): Liao, Cho-Hwa; Pan, Shiow-Lin; Guh, Jih-Hwa; Teng,

Che-Ming

CORPORATE SOURCE: College of Medicine, Pharmacological Institute,

National Taiwan University, Taipei, 100, Taiwan
Biochemical Pharmacology (2004), 67(11), 2031-2038

SOURCE: Biochemical Pharmacology (2004), 67(11), 2031-2038

CODEN: BCPCA6; ISSN: 0006-2952

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

Genistein, a natural isoflavone phytoestrogen present in soybeans, has been extensively studied as a chemopreventive or therapeutic agent in several types of cancer. The traditional Asian diet is rich in soy products may explain in part why the incidence of breast cancer in Asian women is relatively low. To improve therapeutic benefits, we investigated the combination of genistein with chemotherapeutic agents in phenotypically dissimilar human breast cancer cells, MCF-7 and MDA-MB-231, in which estrogen receptor expression is pos. and neg., resp. In the present study, genistein significantly decreased cell apoptosis induced by tubulin-binding agents, paclitaxel and vincristine. FACScan anal. revealed that genistein also diminished the accumulation of the G2/M phase in the cell cycle caused by tubulin-binding agents. In

situ staining of microtubules revealed that genistein could decrease paclitaxel-induced tubulin polymerization However, in vivo tubulin polymerization assay

revealed that simultaneous treatment of genistein did not change the tubulin/microtubule dynamic. Genistein reduced Bcl-2 phosphorylation triggered by paclitaxel and vincristine without changing Bax protein expression. pls and p21 expression, monitored by Western blocking, was not altered by genistein. However, the expression of cyclin B1 and CDC2 kinase was markedly decreased in combination with genistein. In conclusion, genistein inversely affected tubulin-binding agent-induced apoptosis via down-regulation of cyclin B1/CDC2 kinase expression resulting in reduced Bcl-2 phosphorylation.

REFERENCE COUNT: 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 6 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:310964 CAPLUS

DOCUMENT NUMBER:

140:297495

TITLE:

Synergistic anticancer compositions containing

platinum-isoflavonoids

INVENTOR (S):

Kelly, Graham Edmund

PATENT ASSIGNEE(S):

Novogen Research Pty. Ltd., Australia

SOURCE:

PCT Int. Appl., 63 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PAT	ENT I	NO.			KINI	D :	DATE		I	APPL	ICAT:	ION I	NO.		D	ATE		
							-												
	WO	2004	0306	62		A1		2004	0415	Ţ	NO 2	003-2	AU12	96		20	0031	002	
		W:	AE,	AG,	ĀL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	ΒY,	ΒZ,	CA,	CH,	CN,	
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	GE,	
			GH,	GM,	HR,	HU,	ID,	ΙL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	
			LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NI,	NO,	NZ,	
	OM, PG, PH						PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	TJ,	TM,	
	TN, TR, TT						UA,	UG,	US,	UZ,	VC,	VN,	YÜ,	ZA,	ZM,	ZW			
	RW: GH, GM, KE						MW,	MZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,	
	KG, KZ, MD						TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	
				,	,	•		IE,		•	•	•	•	•					
BF, BJ, CF						CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NΕ,	SN,	TD,	TG	
PRIO	RITY	APP	LN.	INFO	.:					i	AU 2	002-	9518	33	ì	4 2	0021	002	
			/ ~ \						~ ~										

OTHER SOURCE(S): MARPAT 140:297495

AB This invention relates to combination therapies involving anticancer chemotherapeutic agents and isoflavones or

analogs thereof. The invention further relates to compds., compns., methods and therapeutic uses involving, containing, comprising, including and/or for preparing platinum-isoflavonoid complexes suitable for use in the combination therapies of the invention. The effect of a composition comprising cisplatin and dehydroequol on various cancer cell lines in culture plates was assessed. It was found that the amount of cisplatin needed to kill a set number of cancer cells was less when in admixt. with dehydroequol as compared to controls with cisplatin alone. Dehydroequol was found to exhibit a strong synergistic interaction with cisplatin in cell lines derived from ovarian, prostate and pancreatic cancers.

REFERENCE COUNT:

THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 7 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2003:965772 CAPLUS

DOCUMENT NUMBER:

140:385592

TITLE:

Cytostatic and cytotoxic activity of synthetic genistein glycosides against human cancer cell lines

AUTHOR(S): Polkowski, Krzysztof; Popiolkiewicz, Joanna;

Krzeczynski, Piotr; Ramza, Jan; Pucko, Wieslaw; Zegrocka-Stendel, Oliwia; Boryski, Jerzy; Skierski, Janusz S.; Mazurek, Aleksander P.; Grynkiewicz,

CORPORATE SOURCE: National Institute of Public Health, Warsaw, 00-725,

SUURCE. Cancer Letters (Amsterdam, Netherlands; (250%),

203(1), 59-69

CODEN: CALEDO; ISSN: 0304-3835

Elsevier PUBLISHER: Journal DOCUMENT TYPE: LANGUAGE: English

Genistein, the principal soy isoflavone, is a mol. of great interest as an innovative chemotherapeutic agent or as a lead-compound in anticancer drug design. To enhance intrinsic activity of genistein and to explore its pharmacophoric potential, its glycosidic derivs. were synthesized. On the basis of structural features and calculated lipophilicity coefficient (C log P) the derivs. were classified as hydrophilic (i.e. those containing free sugar moiety) or lipophilic (i.e. those with alkylated or acylated sugar hydroxyls). The in vitro cytostatic and cytotoxic studies showed hydrophilic glycosides to be practically inactive against human cancer cell lines when compared to the free aglycon. On the contrary, lipophilic glycosides were significantly more active than the parent isoflavone although the correlation between C log P and the activity was not clear. On the basis of GI50 and LC50 values two of the most active glycosides were found to be several times more potent in their cytostatic and cytotoxic effect than genistein. Addnl. all lipophilic glycosides were revealed to exhibit different mode of action in comparison to genistein. It may suggest that these compds. do not undergo

biol. effects primarily as intact mols. REFERENCE COUNT: THERE ARE 31 CITED REFERENCES AVAILABLE FOR THIS 31 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

rapid biodegrdn., either in culture media or inside cells, and exert their

L13 ANSWER 8 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

2003:376630 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: . 138:374200

Chemoprotectant compositions containing TITLE:

> isoflavones Shapiro, Alla

INVENTOR(S):

PATENT ASSIGNEE(S): USA

SOURCE: PCT Int. Appl., 23 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	ENT	NO.			KIN	D	DATE			APPL	ICAT	ION 1	NO.		D	ATE	
WO	2003	0395	37		A1	_	2003	0515	1	WO 2	002-	US35	437		2	0021	105
	W:	ΑE,	AG,	AL.	ΔM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	ВY,	BZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KG,	ΚP,	KR,	KZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	ΝZ,	OM,	PH,
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	US,	UZ,	VN,	ΥU,	ZA,	ZM,	ZW,	AM,	ΑZ,	BY,	KG,	KZ,	MD,	RU,
		TJ,	TM														
	RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ,	ΤZ,	UG,	ZM,	ZW,	ΑT,	BE,	BG,
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		PT,	SE,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,
		ΝE,	SN,	TD,	TG												
DRITY	APP	LN.	INFO	. :					•	US 2	001-	3309	76P		P 2	0011	105

OTHER SOURCE(S):

MARPAT 138:374200

A non-toxic and effective isoflavone chemoprotectant agent for

treating or preventing effects and damage due to the administration of chemotherapeutic agents in the treatment of cancer and other conditions and diseases is described. The isoflavone can be administered orally, s.c., i.m., i.v., transdermally, intranasally, or rectally. The isoflavone is administered chronically, and/or before, during and/or after administration of the chemotherapeutic agent. For example, in pacients with breast cancer undergoing treatment with chemotherapeutic agents that cause severe cardiac toxicity, administration of genistein (0.1-1000 mg/kg) prior and during chemotherapy resulted in decreased cardiotoxicity, allowing an increase in drug intensity, shortened delay in drug administration between doses of the chemotherapeutic

agent, and reduced side effects.

REFERENCE COUNT: THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS 1 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 9 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2003:261608 CAPLUS

DOCUMENT NUMBER:

138:265631

TITLE:

Methods and compositions to determine the

chemosensitizing dose of suramin used in combination

INVENTOR(S):

Au, Jessie L.-S.; Wientjes, M. Guillaume

PATENT ASSIGNEE(S): USA

SOURCE:

PCT Int. Appl., 47 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PAT	CENT 1	NO.					DATE						NO.			ATE	
						-									-		
WO	2003	0265	74		A2	-	2003	0403	1	WO 2	002-1	JS30:	210		2	0020	924
WO	2003	0265	74		Α3		2004	0415									
	W:	ΑE,	AG,	AL,	AU,	BA,	BB,	BG,	BR,	BZ,	CA,	CN,	CO,	CR,	CU,	CZ,	DM,
		DZ,	EC,	EE,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	ΚE,	KΡ,
		KR,	KZ,	LC,	LK,	LR,	LS,	LT,	LV,	MA,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,
		NZ,	OM,	PH,	PL,	RO,	SD,	SG,	SI,	SK,	SL,	TN,	TT,	TZ,	UA,	UG,	US,
		UZ,	VΝ,	YU,	ZA,	ZM,	ZW										
	RW: GH, GM, KE						ΜZ,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	ΑM,	ΑZ,	BY,
		KG,	ΚZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,
		FI,	FR,	GB,	GR,	ΙE,	IT,	ĽU,	MC,	NL,	PT,	SE,	SK,	TR,	BF,	ВJ,	CF,
		CG,	CI,	CM,	GΑ,	GN,	GQ,	GW,	ML,	MR,	ΝE,	SN,	TD,	TG			
EP	1429	713			A2		2004	0623		EP 2	002-	7663	46		2	0020	924
	R: AT, BE, CH						ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
	IE, SI, LT						RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	SK		
US	2004	1809	55		A1		2004	0916	1	US 2	004-	8076	20		2	0040	324
PRIORITY	Y APP	LN.	INFO	.:					1	US 2	001-	3247	04P]	P 2	0010	924
									1	WO 2	002-1	US30:	210	Ţ	W 2	0020	924

A method for determining a therapeutically effective amount of suramin for administering to a patient who is to receive a cytotoxic agent comprises determining the circulating suramin concentration in the patient; administering suramin, if required, to establish a low circulating concentration of suramin in

the patient of below about 200 $\mu\text{M};$ and administering the chemotherapeutic agent to the patient when the low circulating concentration of suramin is present in the patient. Conveniently a nomogram can be constructed for use in clin. settings with the suramin.

L13 ANSWER 10 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:695764 CAPLUS

DOCUMENT NUMBER:

137:210932

TITLE:

Combination therapy for reduction of toxicity of

chemotherapeutic agents

INVENTOR(S):

Prendergast, Patrick T.

PATENT ASSIGNEE(S):

Ire.

SOURCE:

PCT Int. Appl., 66 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NOW. COUNT:

PATENT THEORMATION:

	PAT	CENT	NO.			KIN	D	DATE		i	APPL	ICAT	ION I	. 01		D	ATE	
	WO	2002	0699	49		A2	-	2002	0912	Ţ	WO 2	002 <i>-</i> :	IB63	2		20	0020	305
	WO	2002	0699	49		A3		2003	0605									
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KΕ,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
			PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,
			UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW						•	
		RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SĿ,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,
			KG,	KΖ,	MD,	RU,	ΤIJ,	Tid,	AT,	BE,	CH,	CY,	DE,	DK,	ES,	Fì,	FR,	GB,
			GR,	ΙE,	IT,	LU,	MC,	NL,	PT,	SE,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,
			GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG							
	US	2002	1691	40		A 1		2002	1114	1	US 2	002-	9185	5		2	0020	306
PRIC	RIT	Y APP	LN.	INFO	. :						IE 2	001,-	209		1	A 20	0010	306
AB	Pro	ovide	d in	the	pre	sent	inv	enti	on a	re c	pdwo	s. s	uital	ble :	for	treat	ting	
	nec	oplas	ms a	nd t	umor	s, v	iral	, ba	cter.	ial	and	para	site	inf	ecti	ons a	and	
	cor	nbina	tion	the	rapy	wit:	h th	ese	agen'	ts t	o lo	wer	the a	adve:	rse :	side	eff	ects.

L13 ANSWER 11 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:512399 CAPLUS

DOCUMENT NUMBER:

138:214709

TITLE:

5,6-Dimethylxanthenone-4-Acetic Acid (DMXAA): a New Biological Response Modifier for Cancer Therapy Zhou, Shufeng; Kestell, Philip; Baguley, Bruce C.;

AUTHOR(S):

Paxton, James W.

CORPORATE SOURCE:

Division of Pharmacology and Clinical Pharmacology, Faculty of Medical and Health Sciences, The University

of Auckland, Auckland, N. Z.

SOURCE:

Investigational New Drugs (2002), 20(3), 281-295

CODEN: INNDDK; ISSN: 0167-6997 Kluwer Academic Publishers

PUBLISHER: DOCUMENT TYPE:

Journal; General Review

LANGUAGE: English

A review. The investigational anti-cancer drug 5,6-dimethylxanthenone-4acetic acid (DMXAA) was developed by the Auckland Cancer Society Research Center (ACSRC). It has recently completed Phase I trials in New Zealand and UK under the direction of the Cancer Research Campaign's Phase I/II Clin. Trials Committee. As a biol. response modifier, pharmacol. and toxicol. properties of DMXAA are remarkably different from most conventional chemotherapeutic agents. Induction of cytokines (particularly tumor pecrosis factor (TNF- α), serotopin and nitric oxide (NO)), anti-vascular and anti-angiogenic effects are considered to be major mechanisms of action based on in vitro and animal studies. In cancer patients of Phase I study, DMXAA also exhibited various biol. effects, including induction of $TNF-\alpha$, serotonin and NO, which are consistent with those effects observed in in vitro and animal studies. Preclin. studies indicated that DMXAA had more potent anti-tumor activity compared to flavone-8-acetic acid (FAA). In contrast to FAA that did not show anti-tumor activity in cancer patients, DMXAA (22 mq/kg by i.v. infusion over 20 min) resulted in partial response in one patient with metastatic cervical squamous carcinoma in a Phase I study where 65 cancer patients were enrolled in New Zealand. The maximum tolerated dose (MTD) in mouse, rabbit, rat and human was 30, 99, 330, and 99 mg/kg resp. The dose-limiting toxicity of DMXAA in cancer patients included

acute reversible tremor, cognitive impairment, visual disturbance, dyspnoea and anxiety. The plasma protein binding and distribution into blood cells of DMXAA are dependent on species and drug concentration DMXAA is extensively metabolized, mainly by glucuronidation of its acetic acid side chain and 6-methylhydroxylation, giving rise to DMXAA acyl glucuronide (DMXAA-G), and 6-hydroxymethyl-5-methylxanthenone-4-acetic acid (6 Dil-WAAA), which are excreted into bile and trine. blocks G has been shown to be chemical reactive, undergoing hydrolysis, intramol, migration and covalent binding. Studies have indicated that DMXAA glucuronidation is catalyzed by uridine diphosphate glucuronosyltransferases (UGT1A9 and UGT2B7), and 6-methylhydroxylation by cytochrome P 450 (CYP1A2). Non-linear plasma pharmacokinetics of DMXAA has been observed in animals and patients, presumably due to saturation of the elimination process and plasma protein binding. Species differences in DMXAA plasma pharmacokinetics have been observed, with the rabbit having the greatest plasma clearance, followed by the human, rat and mouse. In vivo disposition studies in these species did not provide an explanation for the differences in MTD. Co-administration of DMXAA with other drugs has been shown to result in enhanced anti-tumor activity and alterations in pharmacokinetics, as reported for the combination of DMXAA with melphalan, thalidomide, cyproheptadine, and the bioreductive agent tirapazamine, in mouse models. Species-dependent DMXAA-thalidomide pharmacokinetic interactions have been observed Co-administration of thalidomide significantly increased the plasma area of the plasma concentration-time curve (AUC) of DMXAA in mice, but had no effect on DMXAA's pharmacokinetics in the rat. It appears that the pharmacol. and toxicol. properties of DMXAA as a new biol. response modifier are unlikely to be predicted based on preclin. studies. Similar to many biol. response modifiers, DMXAA alone did not show striking anti-tumor activity in patients. However, preclin. studies of DMXAA-drug combinations indicate that DMXAA may have a potential role in cancer treatment when co-administered with other drugs. Further studies are required to explore the mol. targets of DMXAA and mechanisms for the interactions with other drugs co-administered during combination treatment, which may allow for the optimization of DMXAA-based chemotherapy.

REFERENCE COUNT:

75 THERE ARE 75 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 12 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2002:190263 CAPLUS

TITLE:

Synthesis, SAR and biological evaluation of zapotin as

a chemoprevention agent

AUTHOR (S):

Hirschelman, Wendy H.; Park, Eun Jung; Murillo, Genoveva; Hawthorne, Michael; Kosmeder, Jerome W., II;

Mehta, Rajendra; Pezzuto, John M.; Moriarty, Robert M.

CORPORATE SOURCE:

DOCUMENT TYPE:

Department of Chemistry, College of Liberal Arts and Sciences, University of Illinois at Chicago, Chicago,

IL, 60607, USA

SOURCE:

Abstracts of Papers, 223rd ACS National Meeting, Orlando, FL, United States, April 7-11, 2002 (2002), MEDI-144. American Chemical Society: Washington, D.

С.

CODEN: 69CKQP

Conference; Meeting Abstract

LANGUAGE: English

AB Zapotin (5,6,2',6,-tetramethoxyflavone, 13) is a natural product found in the seeds of the edible plant, Casimiroa edulis. This flavone was isolated and evaluated in HT-29 and MMOC assays and found to be active as a chemopreventive and chemotherapeutic agent. Based on these results, zapotin (13) and other structurally modified analogs 1-12, 14 were synthesized. The synthetic samples were tested in HL-60 (leukemia cell line), MMOC (mouse mammary organ culture), HT-29 (colon cancer carcinoma cell line) and QR (quinone reductase) screening assays. A structure-activity (SAR) model was established and three potent chemoprevention agents were discovered.

L13 ANSWER 13 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2001:833064 CAPLUS

DOCUMENT NUMBER:

135:352781

TITLE:

Compositions and methods for protecting cells during

cancer chemotherapy and radiotherapy

INVENTOR (S).

Full, William & .; Raghavachari, Malimi; Shu, Hing;

Kink, John

PATENT ASSIGNEE(S):

Wisconsin Alumni Research Foundation, USA

SOURCE:

PCT Int. Appl., 75 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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PATENT NO.
                      KIND DATE
                                       APPLICATION NO.
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                   A1 20011115 WO 2001-US14464
    WO 2001085142
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
           CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
           HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
           LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
           SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
           YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
           DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
            BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                       AA
                             20011115 CA 2001-2408152
    CA 2408152
                             20030205
                                       EP 2001-933017
    EP 1280556
                       A1
          AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
            IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                   T2 20040527
                                      JP 2001-581796
    US 2005043224
                       A1
                             20050224
                                        US 2004-881028
                                                              20040630
PRIORITY APPLN. INFO.:
                                         US 2000-565714
                                                          A 20000505
                                         WO 2001-US14464 W 20010504
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Compns., pharmaceutical prepns. and methods are disclosed for protecting AB non-neoplastic cells from damage caused by cancer chemotherapeutic agents or radiation therapy, during the course of cancer therapy or bone marrow transplant. These are based on the use of chemoprotective inducing agents that induce or increase production of cellular detoxification enzymes in target cell populations. The compns. and methods are useful to reduce or prevent hair loss, gastrointestinal distress and lesions of the skin and oral mucosa that commonly occur in patients undergoing cancer therapy. Also disclosed is a novel assay system for identifying new chemoprotective inducing agents.

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 14 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2000:759656 CAPLUS

DOCUMENT NUMBER:

134:13161

TITLE:

Polymethoxyflavonoids from Vitex rotundifolia inhibit proliferation by inducing apoptosis in human myeloid

leukemia cells

AUTHOR (S):

Ko, W. G.; Kang, T. H.; Lee, S. J.; Kim, N. Y.; Kim,

Y. C.; Sohn, D. H.; Lee, B. H.

CORPORATE SOURCE:

College of Pharmacy and Medicinal Resource Research Center, Wonkwang University, Chonbuk, 570-749, S.

Korea

SOURCE:

Food and Chemical Toxicology (2000), 38(10), 861-865

CODEN: FCTOD7; ISSN: 0278-6915

PUBLISHER:

Elsevier Science Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

AB Three polymethoxyflavonoids from the fruit of Vitex rotundifolia, namely 2',3',5-trihydroxy-3,6,7-trimethoxyflavone (Vx-1), vitexicarpin (Vx-5) and artemetin (Vx-6), were tested for their antiproliferative activity in human myeloid leukemia HL-60 cells. They showed a dose-dependent decrease in the growth of HL-60 cells. The concns. required for 50% inhibition of the growth (IC50) after 96 h were 4.03 μM , 0.12 μM and 30.08 μM for νA 1, V A+5 and V A+6, resp. Treatment of HL-60 cells with the flavonoids induced morphol. changes that are characteristic of apoptosis. We judged the induction of apoptosis by the detection of DNA fragmentation in agarose gel electrophoresis and the degree of apoptosis was quantified by a double-antibody sandwich ELISA and by flow cytometric anal. The C-3 hydroxyl and C-8 methoxyl groups were found not to be essential for the activity, but the C-3' methoxyl instead of hydroxyl group lowered the antiproliferative and apoptosis inducing activity. These results suggest that the polymethoxyflavonoids isolated from V. rotundifolia may be used as potential chemopreventive and chemotherapeutic agents.

REFERENCE COUNT:

THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

LI3 ANSWER 15 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

18

ACCESSION NUMBER:

2000:646580 CAPLUS

DOCUMENT NUMBER:

133:317103

TITLE:

SOURCE:

The therapeutic potential of flavonoids

AUTHOR(S): Wang, Hui-Kang

CORPORATE SOURCE:

University of North Carolina, Chapel Hill, NC, USA Expert Opinion on Investigational Drugs (2000), 9(9),

2103-2119

CODEN: EOIDER; ISSN: 1354-3784

PUBLISHER:
DOCUMENT TYPE:

Ashley Publications Ltd. Journal; General Review

LANGUAGE: English

AB A review with 92 refs. Four most widely investigated flavonoids, flavopiridol, catechins, genistein and quercetin are reviewed in this article. Flavopiridol is a novel semisynthetic flavone analog of rohitukine, a leading anticancer compound from an Indian tree. Flavopiridol inhibits most cyclin-dependent kinases and displays unique anticancer properties. It is the first cyclin-dependent kinase inhibitor to be tested in Phase II clin. trials. Catechin and its gallate are major ingredients in green tea and their anti-oxidant and cancer preventive effects have been widely investigated. A Phase I study of green tea extract GTE-TP91 has been conducted in adult patients with solid tumors. Similarly, genistein is a major ingredient in soybean and has been shown to prevent cancer and have antitumor, anti-oxidant and

anti-inflammatory effects. Two antibody-genistein conjugates, B43-genistein and EGF-genistein, are currently in clin. development for the treatment of acute lymphoblastic leukemia and breast cancer, resp. Finally, most recent updates of quercetin are briefly described.

REFERENCE COUNT:

THERE ARE 92 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

U-13 ANSWER 16 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2000:296079 CAPLUS

DOCUMENT NUMBER:

133:4163

TITLE:

Dietary bioflavonoids induce cleavage in the MLL gene

and may contribute to infant leukemia

AUTHOR (S):

Strick, Reiner; Strissel, Pamela L.; Borgers, Susanne;

Smith, Steve L.; Rowley, Janet D.

CORPORATE SOURCE:

Department of Medicine, Section of

Hematology/Oncology, University of Chicago, Chicago,

IL, 60637, USA

SOURCE:

Proceedings of the National Academy of Sciences of the

United States of America (2000), 97(9), 4790-4795

CODEN: PNASA6; ISSN: 0027-8424

PUBLISHER:

National Academy of Sciences

DOCUMENT TYPE: Journal LANGUAGE: English

Chromosomal translocations involving the MLL gene occur in .apprx.80% of infant leukemia. The search for possible agents inducing infant leukemia identified bioflavonoids, natural substances in food and dietary supplements, that cause site-specific DNA cleavage in the MLL breakpoint cluster region (bCR) in vivo. The MLD BCR DNA cleavage was shown in primary progenitor hematopoietic cells from healthy newborns and adults and in cell lines; it colocalized with the MLL BCR cleavage site induced by chemotherapeutic agents, such as etoposide (VP16) and doxorubicin (Dox). Both in vivo and addnl. in vitro expts. demonstrated topoisomerase II (topo II) as the target of bioflavonoids similar to VP16 and Dox. Based on 20 bioflavonoids tested, we identified a common structure essential for the topo II-induced DNA cleavage. Reversibility expts. demonstrated a religation of the bioflavonoid and the VP16-induced MLL cleavage site. The observations support a 2-stage model of cellular processing of topo II inhibitors. The first and reversible stage of topo II-induced DNA cleavage results in DNA repair, but also rarely in chromosome translocations, whereas the second nonreversible stage leads to cell death because of accumulated DNA damage. Thus, maternal ingestion of bioflavonoids may induce MLL breaks and potentially translocations in utero leading to infant and early childhood leukemia. REFERENCE COUNT: 46 THERE ARE 46 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 17 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

KIND

ACCESSION NUMBER: 2000:227537 CAPLUS

DOCUMENT NUMBER: 132:262172

TITLE: Use of neoangiogenesis markers for diagnosis and

treatment of tumors

INVENTOR(S): Krause, Werner; Muschick, Peter PATENT ASSIGNEE(S): Schering Aktiengesellschaft, Germany

SOURCE: PCT Int. Appl., 27 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.

							-									_			
	WO 2000018439					A2 20000406					WO 1999-EP7198						19990929		
	WO 2000018439 A3 20000914																		
		W:	ΑE,	AL,	AM,	AU,	AΖ,	BA,	BB,	BG,	BR,	BY,	CA,	CN,	CR,	CU,	CZ,	DM,	
•								GM,											
			KZ,	LC,	LK,	LR,	LS,	LT,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	
								sī,											
				YU,															
		RW:	AT,	BE,	CH,	CY,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	
			PT,																
	DE	1984	5798			A 1		2000	0413		DE 1:	998-	1984	5798		1	9980	929	
PRIORITY APPLN. INFO.: DE 1998-19845798 A 19980929										929									
AB	AB Neoangiogenesis markers (i.e. antibodies or receptors for e.g. vascular											ular							
	endothelial growth factor, placenta growth factor, acidic or basic FGF,																		
	transforming growth factor α or β , hepatocyte growth factor,																		
	ins	sulin	-lik	e gr	owth	fac	tor	I, g	lyco	prot	ein 1	B61,	pro	tein	LER	K-1,	flk	-1	
	rec	cepto	r, e	tc.)	or	part	ial	seque	ence	s th	ereo	f and	d and	tian	giog	enic	com	ods.	
	and	l fac	tors	suc	h as	pac	lita	xeĺ,	ende	osta	tin,	fib	rone	ctin	pep	tide	, and	d.	
	fur	nagil	lin a	are d	conj	ugat	ed w	ith a	acti	ve a	gent:	s su	ch a	s				•	
	che	emoth	erap	euti	c ag	ents	, ra	dios	ensi	tize	rs,								
	pho	otose:	nsit:	izer	s, a	ntib	odie	s, o	ligo	nucl	eoti	des,	rad:	ioac	tive	met	al		
	cor	nplex	es, e	etc.	, wh	ich 1	may 1	be bo	ound	to	carr	iers	, fo	r tr	eatm	ent (of t	umors.	
	Lil	cewis	e, n	eoan	giog	enes.	is m	arke	rs m	ay b	e co	njuga	ated	to	diag	nost.	ic a	gents	
		ch as																	
	dia	agnos	is.	Thu	s, N	', N'	, N ' '	', N'	''-t	etra	kis(tert	-but	охус	arbo	xyme	thyl) -N''-	

DATE APPLICATION NO.

DATE

(hydroxycarboxymethyl)diethylenetriamine was converted to its N-hydroxysuccinimide ester, coupled to a Thy-1 antibody, complexed with 186Re, and injected i.v. into rabbits for detection of implanted VX2 tumors by scintigraphy with a gamma camera.

L13 ANSWER 18 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:144761 CAPLUS

<u> росимемт иимъер:</u> 133-103253

TITLE: Activation and protection of T-cells (CD4+ and CD8+)

using an H2 receptor agonist and other T-cell

activating agents

INVENTOR(S): Hellstrand, Kristoffer; Hermodsson, Svante; Gehlsen,

Kurt R.

PATENT ASSIGNEE(S): Maxim Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 38 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

										APPLICATION NO.									
	WO 2000010600								WO 1999-US19211										
WO	WO 2000010600					A3 20000615													
	W:	ΑE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,		
		CZ,	CZ,	DE,	DE,	DK,	DK,	DM,	EE,	EE,	ES,	FI,	FI,	GB,	GD,	GE,	GH,		
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,		
		LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,		
		SD,	SE,	SG,	SI,	SK,	SK,	SL,	TJ,	TM,	TR,	TT,	UA,	UG,	UZ,	VN,	YU,		
		ZA,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM							
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,	DK,		
		ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,		
		CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG							
CA	CA 2341742				AA 20000302			CA 1999-2341742					19990824						
EP								EP 1999-943853											
	R:	AT,	ΒE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,		
		IE,	SI,	LT,	LV,	FI,	RO												
JР	JP 2002523378					T2 20020730			JP 2000-565920					19990824					
TW	TW 576745					20040221				TW 1999-88114376					19990922				
AU	AU 9956870					20000314				AU 1999-56870					19990924				
AU	B2		2003	0925				•											
ZA	A		2001	0927		ZA 2	001-	1787			2	0010	302						
US 2003039628							2003	0227	•	US 2	002-	2655	21		2	0021	003		
PRIORITY APPLN. INFO.:										US 1	998-	1392	81	1	A 1	9980	824		
									1	WO 1	.999-1	US19	211	1	W 1	9990	824		

AB The present invention relates to a method for facilitating activation of T-cells in a patient, comprising: identifying a patient in need of enhanced T-cell activity, administering an effective amount of a T-cell activating composition to the patient, and administering an effective amount

compound that inhibits the production or release of intercellular reactive oxygen metabolites (ROM) to the patient. The present invention further relates to the use of H2-receptor agonists to augment the effectiveness of vaccines. The vaccine composition may also comprises chemotherapeutic agent and/or antiviral agent.

L13 ANSWER 19 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1999:574904 CAPLUS

DOCUMENT NUMBER: 131:295069

TITLE: Oxidative metabolism of monensin in rat liver

microsomes and interactions with tiamulin and other

chemotherapeutic agents: evidence

for the involvement of cytochrome P-450 3A subfamily

AUTHOR(S): Nebbia, Carlo; Ceppa, Luciano; Dacasto, Mauro;

Carletti, Monica; Nachtmann, Carlo

CORPORATE SOURCE: Department of Animal Pathology, Division of

Pharmacology and Toxicology, University of Turin,

Turin, 10126, Italy

SOURCE: Drug Metabolism and Disposition (1999), 27(9),

1039-1044

CODAN: DHDSAI; 188W: 0090-9886

DIBI.ISHER: American Society for Pharmacology and Experimental

Therapeutics

DOCUMENT TYPE:

Journal

English LANGUAGE:

Monensin (MON) is an ionophore antibiotic widely used in veterinary practice as a coccidiostatic or a growth promoter. The aims of this study were to characterize the P 450 isoenzyme(s) involved in the biotransformation of the ionophore and to investigate how this process may be affected by tiamulin and other chemotherapeutic agents known to produce toxic interactions with MON when administered concurrently in vivo. In liver microsomes from untreated rats (UT) or from rats pretreated, resp., with ethanol (ETOH), β naphthoflavone (BNAF), phenobarbital (PB), pregnenolone 16α -carbonitrile (PCN), or dexamethasone (DEX), the rate of MON O-demethylation was the following: DEX > PCN > PB » UT = ETOH > (BNAF; similar results were obtained by measuring total MON metabolism In addition, the extent of triacetyloleandomycin-mediated P 450 complexes was greatly reduced by the prior addition of 100 μM MON. In DEX-treated microsomes, MON O-demethylation was found to fit monophasic Michaelis-Menten kinetics (KM = $67.6 \pm 0.01 \mu M$; Vmax = $4.75 \pm$ 0.76 nmol/min/mg protein). Tiamulin markedly inhibited this activity in an apparent competitive manner, with a calculated Ki (Dixon plot) of 8.2 μM and an IC50 of about 25 µM. At the latter concentration, only ketoconazole or metyrapone, which can bind P 450 3A, inhibited MON O-demethylase to a greater extent than tiamulin, whereas α - naphthoflavone, chloramphenicol, or sulphamethasine was less effective. suggest that P 450 3A plays an important role in the oxidative metabolism of MON and that compds. capable of binding or inhibiting this isoenzyme could be expected to give rise to toxic interactions with the ionophore.

REFERENCE COUNT: THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS 36 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 20 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

1998:245794 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 129:36183

TITLE: The protein kinase c (PKC) inhibitor flavopiridol

> (FLAVO) significantly enhances the cytotoxic effect of chemotherapy by promoting apoptosis in gastric cancer

cells

Schwartz, G. K.; Farsi, D.; Greaney, C.; Werner, J.; AUTHOR (S):

Kelsen, D. K.

CORPORATE SOURCE: Department of Medicine, Division of Solid Tumor

Oncology, Laboratory of Gastrointestinal Oncology, Memorial Sloan-Kettering Cancer Center, NY, USA

Progress in Gastric Research 1997, Proceedings of the

International Gastric Cancer Congress, 2nd, Munich,

Apr. 27-30, 1997 (1997), Volume 1, 627-629.

Editor(s): Siewert, Joerg Ruediger; Roder, Juergen D.

Monduzzi Editore: Bologna, Italy.

CODEN: 65WZAO Conference

DOCUMENT TYPE: LANGUAGE: English

SOURCE:

The failure of many chemotherapeutic agents in gastric tumors reflects an inability of these drugs to induce apoptosis. Even with high concns. of paclitaxel and mitomycin-C (MMC), MKN-74 gastric cancer cells lines are resistant to the induction of apoptosis. We are able to show that flavopiridol, a synthetic flavone, significantly enhances the induction of apoptosis by paclitaxel and MMC in

MKN-74 treated cells.

L13 ANSWER 21 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:628243 CAPLUS

DOCUMENT NUMBER: 127:287832

TITLE: Potentiation of apoptosis by flavopiridol in

mitomycin-C-treated gastric and breast canter cells Schwartz, Gary K.; Farsi, Kian; Maslak, Peter; Kelsen,

AUTHOR(S): Schwartz, Gary F.; Farsi, Kian; Masl.
David P.; Spriggs, David

CORPORATE SOURCE: Division of Solid Tumor Oncology, Gastrointestinal

Oncology Research Laboratory, Gastrointestinal Oncology Section, Memorial Sloan-Kettering Cancer

Center, New York, NY, 10021, USA

SOURCE: Clinical Cancer Research (1997), 3(9), 1467-1472

CODEN: CCREF4; ISSN: 1078-0432

PUBLISHER: American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

Flavopiridol (L86-8275) is a synthetic flavone currently undergoing Phase I clin. trials. It is active against a series of human cancer cell lines and has been shown to inhibit a broad range of protein kinases, including cyclindependent kinases and protein kinase C (PKC). Previous studies have shown that the PKC-specific inhibitor safingol significantly enhances the induction of apoptosis by mitomycin-C (MMC) in gastric cancer cells. Because flavopiridol can potentially inhibit PKC, we elected to determine the extent to which flavopiridol would promote MMC-induced apoptosis in both gastric and breast cancer cells. For these studies, MKN-74 gastric cancer cells and MDA-MB-468 breast cancer cells were exposed to either no drug, 1 μg/mL MMC alone, 300 nM flavopiridol alone, or a combination of chemotherapy with flavopiridol for 24 h. Sequence specificity was also examined by first exposing cells to MMC for 24 h followed by flavopiridol for 24 h or to the same drugs in the reverse order. Apoptosis was measured by quant. fluorescence microscopy of nuclear chromatin condensation in cells stained with the dye, bisbenzimide trihydrochloride. Exposure of MKN-74 cells to flavopiridol alone induced apoptosis in 12 ± 1% of the cells, and exposure to MMC alone induced apoptosis in 10 ± 1%. However, the combination of flavopiridol and MMC increased the induction of apoptosis to 55 \pm 3% of the cells (P < 0.005 for the drug combination vs. flavopiridol alone). Pretreatment with the PKC activator 3-phorbol 12-myristate 13-acetate only partially reversed this effect (43 \pm 1%; P < 0.025). In MDA-MB-468 cells, flavopiridol alone induced apoptosis in 17 \pm 1% of the cells, and MMC alone induced apoptosis in 10 ± 1% of the cells. The combination of flavopiridol and MMC increased the percentage of MDA-MB-468 cells undergoing apoptosis to 58 \pm 4% (P < 0.005 for the drug combination vs. flavopiridol alone). Sequential treatment with MMC followed by flavopiridol induced apoptosis in 63 ± 2% of the MKN-74 cells (P < 0.05 vs. the concomitant drug combination) and in 76 ± 2% of the MDA-MB-468 cells (P < 0.025 vs. the concomitant drug combination), whereas flavopiridol followed by MMC did not increase the induction of apoptosis in either cell line. As determined by the terminal deoxynucleotidyl transferase labeling of the 3' ends of DNA fragments produced in apoptotic cells, the induction of apoptosis with the combination of flavopiridol and MMC occurred to MKN-74 cells in all phases of the cell cycle (i.e., GO-G1, S, and G2-M). These results indicate that flavopiridol potentiates the cytotoxic effect of the chemotherapeutic agent MMC by promoting drug-induced apoptosis in tumor cells. Sequencing studies suggest that MMC followed by flavopiridol or simultaneous treatment is superior to flavopiridol followed by MMC. The enhancement of MMC-induced apoptosis by flavopiridol may be partially PKC dependent and is not associated with one specific region of the cell cycle.

REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ACCESSION NUMBER: 1997:561488 CAPLUS

DOCUMENT NUMBER: 127:199736

TITLE: Accelerated titration designs for phase i clinical

trials in oncology

AUTHOR(S): Simon, Richard; Freidlin, Boris; Rubinstein, Larry;

Arbuck, Susan G.; Collins, Jerry; Christian, Michaele

€.

COPPOPATE SOURCE: Division of Cancer Treatment, Diagnosis, and Centers,

Cancer Therapy Evaluation Program, National Cancer

Institute, Bethesda, MD, USA

SOURCE: Journal of the National Cancer Institute (1997),

89(15), 1138-1147

CODEN: JNCIEQ; ISSN: 0027-8874

PUBLISHER: Oxford University Press

DOCUMENT TYPE: Journal LANGUAGE: English

Many cancer patients in phase I clin. trials are treated at doses of chemotherapeutic agents that are below the biol. active level, thus reducing their chances for therapeutic benefit. Current phase I trials often take a long time to complete and provide little information about interpatient variability or cumulative toxicity. Our objective was to develop alternative designs for phase I trials so that fewer patients are treated at subtherapeutic dose levels, trials are of reduced duration, and important information (i.e., cumulative toxicity and maximum tolerated dose) needed to plan phase II trials is obtained. We fit a stochastic model to data from 20 phase I trials involving the study of nine different drugs. We then simulated new data from the model with the parameters estimated from the actual trials and evaluated the performance of alternative phase I designs on this simulated data. Four designs were evaluated. Design 1 was a conventional design (similar to the commonly used modified Fibonacci method) using cohorts of three to six patients, with 40% dose-step increments and no intrapatient dose escalation. Designs 2 through 4 included only one patient per cohort until one patient experienced dose-limiting toxic effects or two patients experienced grade 2 toxic effects (during their first course of treatment for designs 2 and 3 or during any course of treatment for design 4). Designs 3 and 4 used 100% dose steps during this initial accelerated phase. After the initial accelerated phase, designs 2 through 4 resorted to standard cohorts of three to six patients, with 40% dose-step increments. Designs 2 through 4 used intrapatient dose escalation if the worst toxicity is grade 0-1 in the previous course for that patient. Only three of the actual trials demonstrated cumulative toxic effects of the chemotherapeutic agents in patients. The average number of patients required for a phase I trial was reduced from 39.9 for design 1 to 24.4, 20.7, and 21.2 for designs 2, 3, and 4, resp. The average number of patients who would be

to have grade 0-1 toxicity as their worst toxicity over three cycles of treatment is 23.3 for design 1, but only 7.9, 3.9, and 4.8 for designs 2, 3, and 4, resp. The average number of patients with grade 3 toxicity as their worst toxicity increases from 5.5 for design 1 to 6.2, 6.8, and 6.2 for designs 2, 3, and 4, resp. The average number of patients with grade 4 toxicity

as their worst toxicity increases from 1.9 for design 1 to 3.0, 4.3, and 3.2 for designs 2, 3; and 4, resp. Accelerated titration (i.e., rapid intrapatient drug dose escalation) designs appear to effectively reduce the number of patients who are undertreated, speed the completion of phase I trials, and provide a substantial increase in the information obtained.

REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 23 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1997:147544 CAPLUS

DOCUMENT NUMBER: 126:180992

expected

TITLE: Flavopiridol (L86-8275): selective antitumor activity in vitro and activity in vivo for prostate carcinoma

cells

AUTHOR(S): Drees, Markus; Dengler, Wolfgang A.; Roth, Thomas;

Labonte, Heike; Mayo, Joseph; Malspeis, Louis; Grever,

Michael; Sausville, Edward A.; Fiebig, Heinz H. Department of Internal Medicine, University of

Freiburg, Freiburg, D-79106, Germany

CODEN: CCREF4; ISSN: 1079-0432

PUBLISHER: American Association for Cancer Research

DOCUMENT TYPE: Journal LANGUAGE: English

CORPORATE SOURCE:

We have selected a panel of human tumor xenografts for in vitro and in vivo studies that allows an indication of selectivity of action of novel chemotherapeutic agents. We report here the antitumor activity of the flavone flavopiridol (previously designated L86-8275), which has been selected for further studies based in part on its behavior in the anticancer drug screening system of the United States National Cancer Institute. Eighteen human tumor and five cell line-derived xenografts established by serial passage in nude mice in our laboratory were used as tumor models for in vitro investigations using a modified double-layer soft agar assay. In vivo investigations were completed in nude mice bearing advanced-stage s.c. growing prostate cancer xenografts. Antitumor activity in vitro (test/control ≤ 30%) of

flavopiridol was observed at the very low concentration of 0.1 $\mathrm{ng/mL}$ in three

four prostatic xenografts and in one melanoma xenograft. Overall, in 14 of 23 (61%) tumor xenografts, drug treatment resulted in a IC70 of <10 ng/mL, demonstrating the high antiproliferative potential of flavopiridol. Toxicity to in vitro bone marrow cultures was evident only at 100 ng/mL, indicating potential high selectivity for susceptible tumor cells. Comparison of tumor cells with bone marrow samples tested showed clear prostate carcinoma and moderate melanoma selectivity. In vivo studies of flavopiridol confirmed antitumor activity in both prostate cancer xenografts investigated. At the maximal tolerated dose of 10 mg/kg/day administered p.o. on days 1-4 and 7-11, flavopiridol effected tumor regression in PRXF1337 and tumor stasis lasting for 4 wk in PRXF1369. We conclude that flavopiridol shows strong prostate- and moderate melanoma-specific antitumor activity in vitro. The prostate antitumor activity is also reflected by the two in vivo models studied. Initial clin. efforts with flavopiridol might consider early evaluation in patients with prostate carcinoma.

L13 ANSWER 24 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1996:590815 CAPLUS

DOCUMENT NUMBER: 125:293021

TITLE: Inhibitors of NO activity for improving therapeutic effectiveness of agents for the treatment of solid

tumors and other disorders

INVENTOR(S): Dewhirst, Mark W.; Meyer, Robert E.; Bonaventura,

Joseph; Deangelo, Joseph

PATENT ASSIGNEE(S): Duke University, USA; Apex Bioscience, Inc.; North

Carolina State University

SOURCE: U.S., 29 pp., Cont.-in-part of U.S. Ser. No. 66, 756.

, CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 5554638	Α	19960910	US 1994-246882	19940520
US 5612310	Α	19970318	US 1993-66756	19930524
CA 2163638	AA	19941208	CA 1994-2163638	19940523
WO 9427585	A1	19941208	WO 1994-US5791	19940523

W: CA, JP RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE EP 705098 19960410 EP 1994-917468 19940523 A1 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE JP 09500366 T2 19970114 JP 1994-500873 19940523 US 5788958 Α 19980804 US 1996-709938 19960906 US 6020308 ri. 20000201 UU 1998 126930 10000731 UC 1003-66756 እን 10030574 PPIORITY APPLN. INFO : US 1994-246882 A 19940520 WO 1994-US5791 W 19940523 US 1996-709938 -A3 19960906

AB The present invention is directed to the use of an inhibitor of NO activity, such as a nitric oxide scavenger or an NO synthase inhibitor, as an antitumor therapy to reduce tumor blood flow and oxygenation. invention is also directed to administration of a nitric oxide scavenger or a nitric oxide synthase inhibitor to enhance the effectiveness of tumor therapy with hypoxic or acidic chemotherapeutic agents or hyperthermia. The invention is also directed to the administration of a nitric oxide synthase substrate to a subject previously administered a nitric oxide synthase inhibitor, in order to selectively inhibit tumor perfusion. In a specific example, administration of cell-free Hb, a nitric oxide scavenger, in conjunction with mitomycin C, a hypoxic cytotoxin, results in a significant delay in tumor growth of a human tumor xenograft in a mouse compared to mitomycin C alone. In another example, the administration of an inhibitor of nitric oxide synthase followed by the administration of a substrate of the enzyme causes a specific irreversible reduction of tumor blood flow, while normal blood flow is restored.

L13 ANSWER 25 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1994:621188 CAPLUS

DOCUMENT NUMBER: 121:221188

TITLE: Induction of multiple cytokine gene expression and

IRF-1 mRNA by flavone acetic acid in a

murine macrophage cell line

AUTHOR(S): Eader, Lou Ann; Gusella, Luca; Dorman, Linda; Young,

Howard A.

CORPORATE SOURCE: Biological Carcinogenesis and Development Program,

Natl. Cancer Inst.-Frederick Cancer Research and Dev.

Center, Frederick, MD, 21702-1201, USA

SOURCE: Cellular Immunology (1994), 157(1), 211-22

CODEN: CLIMB8; ISSN: 0008-8749

DOCUMENT TYPE: Journal LANGUAGE: English

AB Flavone-8-acetic (FAA) acid is a potential

chemotherapeutic agent that has demonstrated strong immunomodulatory activity in murine model systems. The immunomodulatory activity of this drug in murine systems has been linked to its ability to rapidly induce cytokine gene expression in vivo ad in mouse splenocytes ev We have now developed a tissue culture model for studying the mol. basis of induction of cytokine expression by FAA. Using the mouse macrophage cell line, ANA-1, we can demonstrate the direct induction of interferon β (IFN β), interleukin-6 (IL-6), tumor necrosis factor- α (TNF α), and interferon response factor-1 (IRF-1) mRNA expression following treatment with FAA. Furthermore the induction of the IFNB mRNA can occur in the absence of new protein synthesis. Nuclear run-on expts. indicate that at least part of the induction of IFNβ, IL-6, and TNF α mRNA occurs at the transcriptional level while the increase in IRF-1 mRNA appears largely post-transcriptional or due to the production of IFNβ protein. Addnl., expts. using agents that interfere with second messengers demonstrate that activation of the protein kinase C pathway is possibly involved in FAA gene induction. The use of this tissue culture model system should lead to a more complete understanding of the mechanisms involved in FAA-induced gene expression and help determine why this drug is inactive on human cells.

L13 ANSWER 26 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1992:587171 CAPLUS

DOCUMENT NUMBER:

117:187171

TITLE:

AULHOR (S):

Reductase and oxidase activity of rat liver cytochrome

P450 with 2,3,5,6-tetramethylbenzoguinone as substrate

Goepear, Armold R.; Te Roppele, Johan M.; Meve,

Etienne P. A.; Vermeulen, Nico P. E.

CORPORATE SOURCE:

Dep. Pharmacochem., Free Univ., Amsterdam, 1081 HV,

Neth.

SOURCE:

Chemico-Biological Interactions (1992), 83(3), 249-69

CODEN: CBINA8; ISSN: 0009-2797

DOCUMENT TYPE:

English

LANGUAGE:

The main objective of the present study was to investigate the proposed role of cytochrome P 450 in the reductive metabolism of quinones as well as in the formation of reduced oxygen species in liver microsomes from phenobarbital (PB-microsomes) and β- naphthoflavone (BNF-microsomes) pretreated rats. In the present study, 2,3,5,6-tetramethylbenzoquinone (TMQ) was chosen as a model quinone. Anaerobic one-electron reduction of TMQ by PB-microsomes showed relatively strong ESR signals of the oxygen-centered semiguinone free radical (TMSQ), whereas these signals were hardly detectable with BNF-microsomes. Under aerobic conditions TMSQ formation was diminished and concomitant reduction of mol. oxygen occurred in PB-microsomes. Interestingly, TMO-induced superoxide anion radicals, measured by ESR (using the spin trap 5,5'-dimethyl-1-pyrroline-N-oxide), and hydrogen peroxide generation was found to occur with βNF-microsomes as well. Furthermore, SK&F 525-A (a type I ligand inhibitor of cytochrome P 450) inhibited TMO-induced hydrogen peroxide formation in both PB- and BNF-microsomes. However, metyrapone and imidazole (type II ligand inhibitors of cytochrome P 450) inhibited mol. oxygen reduction in BNF-microsomes and not in PB-microsomes. The present study indicates that cytochrome P 450-mediated one-electron reduction of TMQ to TMSQ and subsequent redox cycling of TMSQ with mol. oxygen constitutes the major source for superoxide anion radical and hydrogen peroxide generation in PB-microsomes (i.e. from the reductase activity of cytochrome P 450). However, most of the superoxide anion radical formed upon aerobic incubation of TMQ with BNF-microsomes originates directly from the dioxyanion-ferri-cytochrome P 450 complex (i.e. from the oxidase activity of cytochrome P 450). In conclusion, both the one-electron reduction of TMQ and mol. oxygen were found to be cytochrome P 450 dependent. Apparently, both the reductase and oxidase activities of cytochrome P 450 may be involved in the reductive cytotoxicity of chemotherapeutic

L13 ANSWER 27 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

agents containing the quinoid moiety.

ACCESSION NUMBER:

1991:94723 CAPLUS

DOCUMENT NUMBER:

114:94723

TITLE:

Combination of flavone acetic acid (FAA)

with adriamycin, cis-platinum and

difluoromethylornithine (DFMO) in vitro against human

colon cancer cells

AUTHOR(S):

Neelam, Sarabjit S.; Bernabei, Alvise; Freedland, Curtis; Thompson, Roxanna; Corbett, Thomas H.; Luk,

Gordon D.

CORPORATE SOURCE:

Sch. Med., Wayne State Univ., Detroit, MI, 48201, USA

Investigational New Drugs (1990), 8(3), 263-8

CODEN: INNDDK; ISSN: 0167-6997

DOCUMENT TYPE:

Journal

LANGUAGE:

SOURCE:

English

AB Unresectable solid tumors in the metastatic stage are quite resistant to current chemotherapy and radiation therapy regimens. Flavone acetic acid (FAA) is a novel antitumor agent which appears to work through a different mechanism than the conventional chemotherapeutic

agents. In preclin. studies it has been effective against a variety of transplantable murine and human tumors and appears to be solid tumor-selective. It also has nonoverlapping toxicities as compared to conventional agents. Thus, FAA was examined in vitro against human colon cancer cells and explored whether its effectiveness could be enhanced in combination with other agents such as adriamycin (ADR), cis-platinum (CP), and differential formithine (LTM), an inhibitor of polyamine biosynthesis. Addition of FAA for 24 h in liquid media produced dose-dependent growth inhibition. Using soft agar colony assay, growth was inhibited by 58% by 3 mM FAA and only 1.4% by 0.375 mM FAA. The combination of FAA and cis-platinum produced synergism at the lower doses tested. The combination of FAA and adriamycin produced antagonism at all doses tested and the combination of FAA with DFMO didenot produce results significantly different from DFMO alone. Enhancement of FAA activity can thus be achieved in combination with conventional antitumor agents, but may be drug and dose specific.

L13 ANSWER 28 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

1990:171645 CAPLUS

DOCUMENT NUMBER:

112:171645

TITLE:

Biotransformation of N,N',N"-

triethylenethiophosphoramide: oxidative desulfuration to yield N,N',N"-triethylenephosphoramide associated with suicide inactivation of a phenobarbital-inducible

hepatic P-450 monooxygenase Ng, Sze Fong; Waxman, David J.

CORPORATE SOURCE:

Dana-Farber Cancer Inst., Harvard Med. Sch., Boston,

MA, 02115, USA

SOURCE:

AUTHOR (S):

Cancer Research (1990), 50(3), 464-71

CODEN: CNREA8; ISSN: 0008-5472

DOCUMENT TYPE:

Journal English

LANGUAGE:

LANGUAGE

GI

$$\begin{array}{c|c}
X \\
\parallel \\
N - P - N
\end{array}$$
I, X=S
II, X=O

Oxidative metabolism of the polyfunctional alkylating agent thio-TEPA (I) was studied in isolated rat liver microsomes and purified, reconstituted cytochrome P 450 (P 450) enzyme systems in order to elucidate the pathways of drug oxidation and to identify the possible contributions of individual P 450 enzymes to the bioactivation of this chemotherapeutic agent. Rat liver microsomes were found to catalyze conversion of thio-TEPA to its oxo metabolite, II, in a P 450-dependent reaction that was markedly stimulated by prior in vivo treatment with drug inducers of hepatic P 450 subfamily IIB (phenobarbital), but not by pretreatment with inducers of P 450 subfamilies IA (β - naphthoflavone) or IIE (isoniazid). I depletion and II formation catalyzed by phenobarbital-induced liver microsomes were both inhibited by >90% by antibodies selectively reactive with P 450 PB-4 (gene product IIB1), the major phenobarbital-inducible rat liver microsomal P 450 form, but not by antibodies inhibitory toward 7 other rat hepatic P-450s. Oxidation of I to II was also catalyzed by purified P 450 PB-4 [Km (app) 19 μM; Vmax (app) = 11 mol I metbolized/min/mol P 450 PB-4] following reconstitution of the cytochrome with NADPH P 450 reductase in a lipid environment. Metabolism of I by P 450 PB-4 was associated with a suicide inactivation of the cytochrome characterized by Kinactivation = 0.096/min, KI = 24 µM, and a partition ratio of 136 mol I metabolized/mol P 450 inactivated. The metabolite, however, did not inactivate the cytochrome, nor was it subject

to further detectable metabolism $\,$ In microsomal incubations, metabolism of I $\,$ led

to the inactivation of P 450 PB-4 (steroid 16β -hydroxylase) as well as P 450 IIIA-related enzymes (steroid 6β-hydroxylase) and the P 450-independent enzyme steroid 17β-hydroxysteroid:NADP+ 17-oxidoreductase, as demonstrated by use of the P 450 form-selective scendidal substrute androse - neme-3, 1/-dione. In contrast, little or no inactivation of microsomal P 450 IIA-related enzymes (steroid 7α-hydroxylase) or microsomal NADPH P 450 reductase was observed Substantial protection of steroid 6β-hydroxylase and steroid 17β-hydroxysteroid:NADP+ 17-oxidoreductase but not steroid 16β-hydroxylase (P 450 PB-4) was afforded by the sulfhydryl-containing nucleophiles cysteine and glutathione; this suggests that inactivation of these microsomal enzymes is mediated by diffusible, reactive metabolite(s) of I, but that in the case of P 450 PB-4, inactivation occurs before the metabolite(s) depart from the active site. These findings demonstrate that P 450 PB-4 can oxidize I to chemical reactive metabolite(s) that may potentially contribute to drug cytotoxicity.

L13 ANSWER 29 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1989:68875 CAPLUS

DOCUMENT NUMBER: 110:68875

TITLE: Human embryonic cell growth assay for teratogens with

or without metabolic activation system using

microplate

AUTHOR(S): Tsuchiya, Toshie; Matuoka, Atsuko; Sekita, Setsuko;

Hisano, Takuzo; Takahashi, Atsushi; Ishidate, Motoi,

Jr.

CORPORATE SOURCE: Div. Med. Chem., Natl. Inst. Hyg. Sci., Tokyo, Japan

SOURCE: Teratogenesis, Carcinogenesis, and Mutagenesis (1988),

8(5), 265-72

CODEN: TCMUD8; ISSN: 0270-3211

DOCUMENT TYPE: Journal LANGUAGE: English

In vitro microassay for the screening of teratogens was investigated on cancer chemotherapeutic agents sterigmatocystins and benzimidazoles using human embryonic palatal mesenchymal (HEPM) cells. Five thousand cells were inoculated into each well of 96-well microtiter plates, and cultivated for 24 h, after which the media were changed with new ones that contained various amts. of chems.; after cultivation for an addnl. 72 h, the media were discarded, and cells attached to the tissue plate were fixed and stained with Giemsa's solution; the cell number then was conducted by colony counter. For the metabolic activation, the liver S9 obtained from rats pretreated with phenobarbital and 5,6benzoflavone and cofactors (S9 mix) were added directly to the HEPM cell cultures along with chems. After 6 h, the cultures were exchanged with a fresh medium and incubated for a further 72 h. Concns. of the cancer chemotherapeutic agents that inhibited growth by 50% ranged from 0.001 to 10 µg/mL. Sterigmatocystins indicated strong inhibition; among three derivs., O-acetyl sterigmatocystin was the most potent inhibitor. Benzimidazoles also exhibited an inhibitory action on HEPM cell growth; nitro and chloro groups at the 5 position in 2-(2-pyridyl)benzimidazole were potent substituents. As for the activation of cyclophosphamide in the HEPM cell culture, IC50 was decreased to 1.0 µg/mL by the incubation with S9 mix for 6 h, and sterigmatocystin was activated by S9 mix.

L13 ANSWER 30 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1989:13639 CAPLUS

DOCUMENT NUMBER: 110:13639

TITLE: Dental compositions containing antiphlogistic agents,

antiosteoporotic agents, and local anethetics or

chemotherapeutic agents

INVENTOR(S): Csanyi, Endre; Csanyi, Gabor; Balogh, Tibor; Nagy,

Laszlo

PATENT ASSIGNEE(S): Reanal Finomvegyszergyar, Hung.

SOURCE: PCT Int. Appl., 15 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATINT INFOMMATION:

PA	CENT NO.					DATE		API	PLICATION NO	DATE			
WO	8805650 W: AU, DK, JP,			A1 19880811			0811	WO	1988-HU3		19880129		
	RW: AT	BE,	CH,	DE,	FR,	GB,	IT,	LU, NI	L, SE				
HU	201474			В		1990	1128	HU	1987-360		19870203		
AU	8812263	1		A1		1988	0824	AU	1988-12263		19880129		
EP	349535			A1		1990	0110	EP	1988-901085		19880129		
EP	349535			B1		1992	0108						
	R: AT	BE,	CH,	DE,	FR,	GB,	IT,	LI, L	J, NL, SE				
JP	0250237	'5		T2		1990	0802	JP	1988-501316		19880129		
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CN	8810049	8		Α		1988	0928	CN	1988-100498		19880203		
DD	273977			A5		1989	1206	DD	1988-312633		19880203		
CS	277001			В6		1992	1118	CS	1988-674		19880203		
CA	1327529)		A1		1994	0308	CA	1988-558061		19880203		
ES	2009561			A6		1989	1001	ES	1988-675		19880307		
DK	8805431			Α		1988	0929	DK	1988-5431		19880929		
PRIORIT	Y APPLN.	INFO	. :					HU	1987-360	Α	19870203		
								EP	1988-901085	Α	19880129		
								WO	1988-HU3	Α	19880129		

AB A dental comp., useful as a therapeutic dental filling for preserving teeth during any stage of pulpitis and periodontitis, comprises an antiosteoporotic agent, an antiphlogistic agent, an optional natural or synthetic chemotherapeutic agent or local anesthetic, and ≥1 known auxiliary agents. A mixture containing doxycycline hyclate 2, triamcinolone 0.8, ipriflavone 2.0, ZnO 80, and CaO 15.2 g, was admixed with eugenol to obtain a therapeutic dental cement.

L13 ANSWER 31 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1988:161016 CAPLUS

DOCUMENT NUMBER: 108:161016

TITLE: The structure of flavone-8-acetic acid, a

chemotherapeutic agent, and its application to drug design

AUTHOR(S): Rabinovitz, M.

CORPORATE SOURCE: Div. Cancer Treat., Natl. Cancer Inst., Bethesda, MD,

20892, USA

SOURCE: Journal of Enzyme Inhibition (1988), 2(2), 151-2

CODEN: ENINEG; ISSN: 8755-5093

DOCUMENT TYPE: Journal LANGUAGE: English

AB Pyrone-2-propionic acid may be the active moiety of the resonance form of

flavone-8-acetic acid, an agent with high activity against colon

adenocarcinoma.

L13 ANSWER 32 OF 32 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1949:768 CAPLUS

DOCUMENT NUMBER: 43:768

ORIGINAL REFERENCE NO.: 43:224f-i,225a-i,226a-i

TITLE: Basically substituted xanthone and thiaxanthone

derivatives; miracil, a new chemotherapeutic

agent

AUTHOR(S): Mauss, Hans

SOURCE: Chemische Berichte (1948), 81, 19-31

CODEN: CHBEAM; ISSN: 0009-2940

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

cf. Naturwissenschaften 33, 253(1946). It seemed likely that the preparation of soluble salts of basically-alkylated xanthone derivs., like plasmochin in the quinoline and atebrin in the acridine series, would lead to interesting results. Basically substituted ethers and carboxamides of xanthone (1) showed no chemocherapeuric activity, but replacement of a Cl atom or other reactive substituent adjacent to the CO group by basic residues yielded compds. which in animal expts. gave indication of activity against the causative agents of bilharziasis (schistosomiasis). In what follows, R = 2-diethylaminoethylamino. 1-R-xanthone (II) showed no activity against schistosomes, and it was found that a Me group in the para position to the basic group is essential for activity. Replacement of this Me group by Et, MeO, or even Cl again results in loss of activity, whereas in the atebrin series Cl and Me are mutually replaceable without affecting the activity. Furthermore, in the I series, the activity in general decreased when the number of C atoms in the basic side chain exceeded 2. Starting with a short basic chain and a Me group in the para position to it [1-R-4-methylxanthone, miracil A (IIa)], attempts were made to increase the activity by suitable substitution. This was effected by introduction on the unsubstituted behzene of Me, MeO, or Cl, a 6- proving to be therapeutically more effective than a 7-substituent. In this series 6-chloro-1-R-4-methylxanthone (miracil B) (III) was the most active in the mouse test, but all, except IIa, were inactive (possibly because of poor resorption) against bilharziasis in the ape. Further attempts to obtain more active and more easily resorbable derivs. led to 1-R-4methylxanthydrol (miracil C) (IV), which was more effective both in the mouse and in the ape test. Quaternary basic xanthone salts proved to be inactive. On the other hand, 1-R-4-methylthiaxanthone-HCl (miracil D) (V) was the most effective in the ape test of any compound thus far prepared Variation of the basic group resulted in loss of activity except in the case of the Et2N(CH2)3NH derivative Derivs. of V with a 2nd Me group on the substituted benzene ring or a Cl atom on the other benzene ring retained their activity but replacement of the 4-Me group by MeO or Cl resulted in inactive compds. The sulfone of V was inactive, while the thiaxanthene and its analogs were active. V is to be tested clin. II, yellow crystals from ligroin, m. 99-100°, was obtained from 22.6 g. 1-methoxyxanthone and 35 g. Et2NCH2CH2NH2 heated 5 h. at 190-200°, together with a considerable amount of 1-hydroxyxanthone. o-(5-Chloro-2-methylphenoxy)benzoic acid, m. 117-18° (from alc.), was obtained in almost 90% yield from o-ClC6H4CO2H and 5,2-ClMeC6H3OH according to Ullmann and Zlokasoff [Ber. 38, 2111(1905)]; heated on the water bath with concentrated H2SO4, it gave about 85% 1-chloro-4-methylxanthone (VI), m. 133-4° (from alc.). VI boiled 10 h. in PhNO2 with 1.5 mols. p-MeC6H4SO2NH2, KOAc, and a little Cu bronze, and the resulting 1-sulfonamido compound gently heated 1 h. on the water bath with concentrated H2SO4, yielded about 85% 1-amino-4-methylxanthone (VII), deep yellow crystals from alc., m. 139-40° [Ac derivative, m. 175-6° (from alc.)]. 1-(Methylamino) compound, from VI and alc. MeNH2 heated several hrs. at 180-90°, yellow crystals from benzene-ligroin (1:2), m. 133-4°. 1-(2-Hydroxyethylamino) compound (about 90% from VI and HOCH2CH2NH2 heated several brs. at 170° with pyridine and Cu bronze), yellow, m. 187-8° (from alc.), gives on heating 2 h. on the water bath with excess POC13 about 60% of the 1-(2-chloroethylamino) compound (VIII), yellow needles from alc., m. 145-6°. 1-R compound (IX) (around 90% from VI and Et2NCH2CH2NH2 heated 6 h. at 170°), yellow crystals from alc., m. 76-7°; HCl salt, yellow, m. 190-1°; picrate, reddish yellow crystalline powder from alc., m. 136-7°. IX is also obtained from VII slowly heated (0.5 h.) to 150° with 1.1 mols. Et2NCH2CH2Cl; from VIII heated 6 h. at 190° with excess Et2NH in toluene; and in moderate yield by ring closure of o-(2,5-MeRC6H3O)C6H4CO2H with concentrated H2SO4. 4-R-2-hydroxytoluene, b4 178-9°, obtained by demethylation of the Me ether, b3 154-5°, with concentrated HBr. IX heated 4 h. at 170-80° with alc. KOH, the alc. driven off, water added, and the

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alkaline solution extracted with ether and acidified with dilute HCl yields an
     amphoteric precipitate redissolving in excess of acid; taken up in ether and
    purified through the AcOH salt, it gives 6-R-2,2'-dihydroxy-3-
    methylbenzophenone (X), yellow crystalline powder from alc., m. 88-9°
     [picrate, yellow, m. 177-8° (from alc.)]; during the purification of X
     through the AcOH salt there is obtained, as an acid-insol. byproduct, a
     small amount of I hydroxy-2-methylaanchone, yellow needles from alc., a.
     149-9°, formed in larger yield when Y is boiled a short time in
    AcOH and allowed to cool; the 2 xanthones are formed in the ratio 4:5 in
     this condensation. o-(5-Chloro-2,4-dimethylphenoxy)benzoic acid,
     from o-ClC6H4CO2H and 5,2,4-ClMe2C6H2OH, m. 146-7° [from
    dilute alc. (4:1)], converted by concentrated H2SO4 into 1-chloro-2,4-
    dimethylxanthone, m. 159-60°; 1-R-2,4-dimethylxanthone-HCl, yellow,
    m. 179-80° (from alc.). 4-Chloro-2-(5-chloro-2-
    methylphenoxy)benzoic acid (about 70% from 2,4-Cl2C6H3CO2H and
     5,2-ClMeC6H3OH), m. 176-7° (from alc.); 1,6-dichloro-4-
    methylxanthone (yield almost quant.), m. 177-8° (from glacial
    AcOH), gives with Et2NCH2CH2NH2 the 1-R compound, yellow, m. 87-8°
     (from ligroin) [HCl salt, yellow, m. 255-6° (decomposition);
    methanesulfonate, yellow, m. 142-3°]: 5-Chloro-2-(5-chloro-2-
    methylphenoxy) benzoic acid, yellowish, m. 177-8° (from alc.);
     1,7-dichloro-4-methylxanthone, m. 198° (from glacial AcOH); 1-R
    compound, isolated in about 80% yield as the HCl salt, yellow, m.
     243° (from alc.). 2-(5-Chloro-2-methylphenoxy)-4-methylbenzoic
    acid, m. 138-9° (from alc.); 1-chloro-4,6-dimethylxanthone, needles
    from alc., m. 168°; 1-R compound [about 75% as the HCl salt, yellow,
    m. 217-18° (from alc.)]. 2 - (5 - Chloro - 2 - methylphenoxy) - 5
     - methylbenzoic acid, m. 173-4° (from alc.); 1-chloro-4,7-
    dimethylxanthone, m. 152° (from alc.); 1-R compound [HCl salt,
    yellow, m. 198° (from alc.)]. 2-(5-Chloro-2-methylphenoxy)-4-
    methoxybenzoic acid, m. 174-5° (from alc.); 1-chloro-6-methoxy-4-
    methylxanthone, needles from glacial AcOH, m. 176-7°, greatly
    depresses the m.p. of the 7-MeO derivative below; 1-R compound yellow, m. .
     84-5° (from ligroin), isolated as the yellow HCl salt, m.
    225-6° (decomposition) (from alc.). 2-(5-Chloro-2-methylphenoxy)-5-
    methoxybenzoic acid m. 183° (from alc.); 1-chloro-7-methoxy-4-
    methylxanthone m. 175-6° (from alc.); 1-R compound [HCl salt, yellow,
    m. 189-90° (from alc.)]. IV, from the xanthone in boiling aqueous alc.
    NaOH with Zn dust, needles from ligroin (b. 60-70°), m.
     100°, forms on cautious neutralization with dilute HCl a yellow solution
    which with a slight excess of acid becomes deep blue-green.
    o-HSC6H4CO2H and p-ClC6H4Me with concentrated H2SO4 give a mixture of
     1-chloro-4-methyl-(XI) and 4-chloro-1-methylthiaxanthone (XII) which can
    hardly be separated (Ullmann and v. Glenck, C.A. 11, 2668). Only the Cl atom
    of XI reacts with Et2NCH2CH2NH2, and when 39 g. of the mixture of XI and XII
    was heated 4 h. with an excess (54 q.) of the amine, treated with 100 cc.
    of 2 N NaOH, distilled with steam, the NaOH solution decanted off, the
semisolid
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residue treated with 10% AcOH, and the insol. unchanged XII, m. 141-2°, filtered off, NH4OH or dilute NaOH precipitated from the orange-red filtrate about 22 g. of the free base of V, yellow, m. 64-5° (from alc.); HCl salt (V), yellow, m. 195-6°, soluble in water with orange-yellow color and neutral reaction. The base is also obtained from 1-amino-4-methylthiaxanthone and 1.1 mols. Et2NCH2CH2Cl heated 1 h. at 150°. 1-Acetamido-4-methylthiaxanthone, yellow needles from benzene, m. 180-1°; 1-(2-hydroxyethylamino) compound (about 60% from the mixture of XI and XII with H2NCH2CH2OH), orange needles (from alc.), m. 183-4°, gives with POCl3 the 1-(2-chloroethylamino) compound, yellow, m. 146° (from alc.), converted by Et2NH into V. 1-(3-Diethylaminopropyl) analog of V, from the mixed XI and XII with Et2N(CH2)3NH2 (b16 59-61°), orange-red oil; HCl salt, yellow, m. 1-Chloro-2,4-dimethylthiaxanthone, from o-HSC6H4CO2H and 2,4-Me2C6H3Cl, fallow, m. 143-4° (from glacial AcOH); 1-R compound, orange-yellow, m. 193-4° (from MeOH-Et2O). 1-R-4-methoxythiaxanthone, from the 1-Cl compound, isolated in small yield

as the orange HCl salt, m. 246-8°. 1,6-Dichloro-4methylthiaxanthone (XIII); obtained mixed with the 4,6-dichloro-1-Me isomer (XIV) from 4,2-Cl(HS)C6H3CO2H and p-ClC6H4Me, needles from glacial AcOH, m. 182-3°; 1-R compound, from the mixture of XIII and XIV, red oil soon solidifying and separating from alc. in yellow crystals, m. 96-7° [HCl salt, yellow, m. 246-7° (from glacial AcOH), difficult, soluble in cold, more easily soluble in how water with an orange-vellow color). The residue remaining after extracting the base with dilute AcOH yields from glacial AcOH yellowish crystals of XIV, m. 201°. The mixture (130 q.) of XI and XII stirred 4 h. in 1300 cc. glacial AcOH with 300 cc. of 25% H2O2, let stand overnight, heated 6 h. at 50-60°, and cooled, yields 114 g. of a mixture, m. 177.5°, of the mixed sulfones, which, heated 6 h. at 170-80° with Et2NCH2CH2NH2 and pyridine, gives 1-R-4-methylthiaxanthone 10-dioxide, orange needles from alc., m. 116-17°, isolated as the HCl salt, orange-yellow, m. 230° (from MeOH); the alc. mother liquors from the salt yield the yellow HCl salt, m. 220° (from MeOH), of the 4-R-1-Me dioxide (XV), yellow scales from alc., m. 128-9°. 4-Chloro-1-methylthiaxanthone 10-dioxide (51 g. from 52 g. of the thiaxanthone, m. 141-2°, with H2O2 in glacial AcOH), yellowish white, m. 170-1°, gives XV with Et2NCH2CH2NH2 and Cu bronze in pyridine at 170-5°. 1-R-4-methylthiaxanthene (XVI), from the xanthone in boiling alc. gradually treated with Na, fallow, m. 66-7° (from alc.); HCl salt, m. 142-3°. 1-Amino-4-methylthiaxanthene, prisms from alc., m. 96-7° [Ac derivative, m. 195-6° (from alc.)], gives XVI with Et2NCH2CH2Cl at 150°.

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